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# PROCEEDINGS OF THE ROYAL SOCIETY.

SECTION A — MATHEMATICAL AND PHYSICAL SCIENCES.

## *Discussion on Heavy Hydrogen*

### Opening Address

By LORD RUTHERFORD, O M , F R S

In the history of Physical Science, it is a commonplace that a new discovery which at first appears to be of purely scientific interest, ultimately, within a period of twenty years or more, is found to have useful practical applications. This is well illustrated by the discovery of the rare gases in the atmosphere, neon, and argon, which are now used in quantity for industrial purposes. The fundamental discovery in 1919 of the isotopic constitution of the majority of our elements, so largely due to Aston, at first sight appeared to be of purely scientific significance, but we shall see from the discussion to-day that it may ultimately have wide practical consequences in many directions.

It is hardly necessary to discuss in detail the history of the discovery and separation of heavy hydrogen, in which scientific men in the U S A have taken such a leading part. The proof that oxygen was not a simple element but contained two isotopes in small quantity of masses 17 and 18, indicated that there was a small discrepancy of about 2 parts in 10,000 between the measurements of the relative masses of hydrogen and oxygen found by Aston and those found by direct physical and chemical methods. Birge and Mendel suggested that this discrepancy might be due to the presence of an isotope of mass 2 present in ordinary hydrogen. This gave the necessary impetus to Urey, Brickwedder and Murphy to test whether the presence of  $H^2$  could be detected by direct optical methods. The experiments were successful in showing a small trace of  $H^2$ , estimated initially at about 1 in 4000 of the  $H^1$  isotope. The wave-length of the  $\alpha$  line of  $H^2$  was found to be 1.79 Å units greater than for  $H^1$ —a result agreeing closely with the theoretical value to be expected for an isotope of hydrogen of mass 2. The mass of the new isotope was directly

measured by Bainbridge, using a modified type of mass spectrograph, and found to be 2.0136, slightly less than the weight of the ordinary hydrogen molecule 2.0156 in terms of  $O = 16$

We have no definite evidence of the exact constitution of  $H^3$ , whether it is to be regarded as a simple entity or built up of two or more constituents. It was at first natural to suppose that the  $H^3$  nucleus might be made up of two protons and a negative electron, but the subsequent discovery of the neutron indicated that it might rather be a close combination of a neutron and a proton. Taking Chadwick's value of the mass of the neutron as 1.0067, the sum of the masses of the proton and neutron is 2.0145, while the mass of the  $H^3$  nucleus is slightly less, 2.0136, indicating that the binding energy of the neutron-proton combination is less than 1 million volts. If this be so, it is expected that the  $H^3$  nucleus should be broken up by collision with a swift  $\alpha$ -particle. In conjunction with Mr. Kempton I have made experiments to test this, but have been unable to detect with certainty the presence of any neutrons when heavy water was bombarded by  $\alpha$ -particles from polonium. The number of neutrons, if any, was certainly less than 1% of the number of neutrons released from a sheet of beryllium under the same conditions. If the disruption of  $H^3$  with an emission of a neutron occurs, it must happen very rarely compared with the number of violent collisions between the  $\alpha$ -particles and the  $H^3$  nucleus.

It is interesting to note here a suggestion made by Lawrence. He found in his experiments on the bombardment of matter by high speed  $H^3$  ions that a group of protons of nearly the same speed was released from a number of elements. In explanation, he suggested that the  $H^3$  nucleus broke up into a neutron and proton either in the bombarded nucleus or in the strong field in its neighbourhood. For the conservation of energy to hold, it is necessary to suppose that the mass of the neutron is much lower than that found by Chadwick, viz., 1.0006 instead of 1.0067. On this view, the  $H^3$  nucleus contains a store of energy corresponding to about 5 million volts, and this is occasionally released in nuclear collisions. Further experiments are required to test the validity of this idea.

It was of interest to me also to examine whether the fields of force near the  $H^1$  and  $H^3$  nuclei are the same. This was tested by comparing the distribution with the velocity of the recoil  $H^1$  and  $H^3$  atoms when  $\alpha$ -particles pass through ordinary and heavy hydrogen respectively. While the recoil  $H^3$  particles travel, as is to be expected, slightly further than the  $H^1$  particles, to a first approximation the number and distribution of the recoil atoms were about the same in the two cases. Since in a close collision the  $\alpha$ -particles and the  $H^3$

nucleus approach within  $10^{-13}$  cm. of each other, these results indicate that the scattering fields are sensibly the same for  $H^1$  and  $H^2$  nuclei, even up to these very small distances

Some success in concentrating  $H^2$  was initially obtained by fractionating liquid hydrogen. Washbourn and Urey noted that there was a greater concentration of  $H^2$  in old electrolytic cells and found the  $H^2$  was rapidly enriched in the residues after electrolysis. This general method was first used on a large scale by Lewis and Macdonald, of the University of California, in order to obtain concentrated preparation of heavy water. By this method they have obtained quantities of heavy water of the order of several hundred cubic centimetres practically in a pure state. It has been concluded that one atom of  $H^2$  is normally present with 6500 atoms of ordinary hydrogen. Lewis and his collaborators find that the density of this new water is about 11% higher than that of ordinary water, while its freezing point is  $3.8^\circ C$  and its boiling point  $101.42^\circ C$ . The maximum density is found to occur at  $11.6^\circ C$  instead of  $4^\circ C$  as in normal water. It is of interest also to refer to another method of concentration by pure diffusion methods. Professor Hertz informs me that he has been able to obtain the new isotope in small quantity in a pure state by applying his elaborate diffusion method to ordinary hydrogen. He states that he has obtained heavy hydrogen so pure that he has been unable to detect in its spectrum the  $\alpha$  line of ordinary hydrogen. I shall not discuss further the practical methods of separation, for I hope the methods of obtaining heavy water will be explained to you by Dr. Harteck who has been responsible for a preparation of about 25 c.c. of the new water in the Cavendish Laboratory, for use in experiments on the transformation of matter.

It is obvious that this new discovery opens up a wide and important field of work, but I shall leave my chemical colleagues to deal adequately with this question. On account of its greater mass, it is to be expected that the rate of diffusion and the rate of chemical reaction will differ when  $H^2$  is substituted for  $H^1$ , while the compounds formed with the new isotope are to be expected in some cases to exhibit rather different properties from the normal hydrogen compounds. Similarly, this new discovery opens up interesting questions on the effect of heavy water in altering the normal physical and chemical processes in animal and plant life. A certain amount of information is already available in this interesting field of enquiry.

There is one question of much interest to me to which I should like to refer, namely, the use of  $H^2$  nuclei as swift projectiles for studying the transformation of the elements. It was a happy coincidence that when Professor Lewis had



prepared concentrated samples of  $H^3$ , Professor Lawrence of the same University had in working order his ingenious apparatus for obtaining high-speed ions corresponding to more than a million volts in energy. Lawrence found that the high speed  $H^3$  ions were often much more effective than protons of equal energy in causing transformations of new kinds in a number of elements. For example, when lithium is bombarded with  $H^3$  ions,  $\alpha$ -particles are ejected with speed considerably greater than the swiftest  $\alpha$ -particle from radioactive substances. It is now clear that an  $H^3$  particle occasionally enters the nucleus of lithium mass 6, and the resulting nucleus then breaks up into 2  $\alpha$ -particles, escaping in nearly opposite directions. The correctness of this view is well shown by the Wilson chamber photographs of the tracks of the  $\alpha$ -particles obtained by Dee and Walton. The action of  $H^3$  on the isotope of mass 7 is even more complicated, for Oliphant and I have observed that  $\alpha$ -particles are liberated over a wide range of velocities. In this case, it seems that the capture of  $H^3$  by the lithium nucleus of mass 7 results in the break up of the system into two  $\alpha$ -particles and a neutron. We estimate that the maximum energy of the ejected neutron may be as great as 15 million volts. We have confirmed this conclusion by finding that neutrons can be detected in numbers corresponding to this mode of transformation using  $H^3$  particles of energy about 200,000 volts. Lauritsen found that a copious supply of neutrons could be obtained by bombarding beryllium with  $H^3$  particles, while Lawrence obtained large numbers from lithium with very fast  $H^3$  particles, but he inclines to believe that most of the neutrons observed in his experiments arise from the break up of the  $H^3$  nucleus into a neutron and a proton.

As already mentioned, Lawrence has observed that  $H^3$  bombardment gives rise to one or more groups of fast protons from a number of elements. These observations have been confirmed by Cockcroft and Walton for several light elements such as Li, C and Fe, using  $H^3$  particles of energy about 500,000 volts, but they have failed to observe proton groups from Cu and Au. In general it appears that the  $H^3$  particle is remarkably effective in causing the transformation of many elements, resulting in a number of substances in the liberation of  $\alpha$ -particles as well as protons and neutrons. There can be no doubt that this new projectile, as well as the proton, will prove of great service in studying the processes which take place in the transformation of the elements, and this will give further important information on the structure of nuclei.

It is obvious that this new isotope, which can so easily be obtained in reasonable quantity in a pure state, will prove of such great importance to science that it is desirable to give it a definite name. Urey has proposed the

name "deuterium" for the new isotope. It is important also that an appropriate title should be given to the  $H^2$  nucleus not only as a projectile for atomic transmutations but as a possible constituent of atomic nuclei. Lewis has suggested the name "deuton" or "deuteron" for this nucleus. While we all realize that the first discoverer has a strong claim in suggesting an appropriate name for a new substance the question of a suitable nomenclature is in this case of such general importance to scientific men that it deserves careful consideration. While the name "deuton" is in some ways suitable it has for me the objection that it is liable in the spoken word to be confused with neutron, and this difficulty is accentuated by the recent discovery that neutrons are often liberated from elements bombarded by deutons. In consultation with some of my physical and chemical colleagues some time before these names were announced the name "diplogen" ( $\delta\iota\pi\lambda\omicron\gamma\epsilon\varsigma$ —double) for heavy hydrogen and "diplon" for the nucleus seemed to meet with some favour. Whatever view may be taken on this question it is important that the new isotope should have a definite symbol allotted to it and the symbol "D" seems appropriate.

In this brief statement I have not attempted to deal with a number of interesting points which I hope will receive the attention of subsequent speakers.

**THE PRESIDENT.** I think it might be well if each speaker would express his opinion on this question of the names, it would be useful to have the collective opinions of all those who speak.

**DR N V SIDGWICK, FRS.** I should like, in compliance with the President's wishes, to express very strong support for the terminology suggested by Lord Rutherford. I do not think that the objection he raised to the term "deuteron" because of its similarity to "neutron" is at all an unimportant matter, I have heard from people who were at the meeting of the American Chemical Society at Chicago last summer that they actually had very great difficulty in determining whether a speaker said "neutron" or "deuteron". You must remember that this is not like the discovery of a rare earth. It does not much matter to any of us what we call No. 61 because we rarely have occasion to refer to a substance like that, but the isotope of hydrogen is going to be one of the most important substances in the coming development of chemistry.

The names which Lord Rutherford suggested give us exactly what we want. They give us, first of all, the relation between the nucleus of the new isotope and the nucleus of the old isotope, and it is really more sensible to call it "the

double one " than " the second one " because the interesting thing about it is not that it is the second lightest particle, that is a matter of argument and it may be said that the first particle is the proton and the second the thing with atomic number 2. The interesting thing about it is that it is double the first one, and that is the reason for the name Lord Rutherford suggests. Then you have the relation between the particle and the atom as a whole which is given by "diplogen," which means that which makes diploon, and the only way in which you can make the particle is from the atom of the isotope. "Diplogen," too, conveys the obvious analogy to hydrogen, which is the most striking thing about it. "Deutium," "deuterium" or "deuteronium" do not indicate a relation to hydrogen. Moreover, when we come to compounds like  $\text{CH}_3\text{D}$  and  $\text{C}_6\text{H}_5\text{D}$ , it is quite easy to call them diplomethane or di-diplobenzene, whereas if you call them deuteromethane or di-deutero benzene it must be remembered that the prefix "deutero" is a living prefix in scientific terminology, and means not "containing deuterium," but simply "second."

I have been asked to say something on the chemical side of the question, and I will do so as briefly as I can, since I have no new knowledge to produce. I want to deal with what we may expect to be the properties of substances made of this new material, as compared with those made of hydrogen, and along what lines we may expect the investigation of the new substance to lead to the discovery of new knowledge.

It is to be expected that the atomic volume of the new isotope will be very nearly the same as that of hydrogen, just as the two isotopes of lead have, as far as we know, the same atomic volume. Moreover, since the atomic volume is practically the same and the atomic number is the same, the ordinary physical properties of a substance made of this element will only differ slightly from those of a substance made of ordinary hydrogen. That is the first point.

The second point is that it has been shown by Urey and Rittenberg, on the basis of theoretical physics, that the equilibrium constant for reactions in which the new isotope takes part will not be quite the same as for those in which the old hydrogen takes part. The difference is not likely to be more than 50%, thus for the reaction  $\text{H}_2 + \text{I}_2 = 2\text{HI}$  they find a difference of 20%.

This refers to reactions at the ordinary temperature, if you do them at the absolute zero that difference ought to increase up to infinity, but at the ordinary temperature you may expect rather small differences. With regard to this, we have already the experimental evidence of Lewis that the maximum density of  $\text{D}_2\text{O}$  is at  $11.6^\circ\text{C}$ , whereas that of ordinary water is at  $4^\circ\text{C}$ . We all realize,

of course, that water, at any rate at temperatures near freezing point, is made up of a variety of polymers in some sense or another, and that the abnormal change in density near freezing point is due to that fact. Now, it is obvious from the fact that the maximum density for the  $D_2O$  is at a different temperature from that of water, that the equilibrium between those polymers or between the various states of water is different for the two substances, and therefore—and this, I think, has not always been noticed—it follows that the differences which have been observed in the physical properties of the heavy water and the light water—the difference of  $3.8^\circ$  in freezing point and  $1.4^\circ$  in boiling point, about  $\frac{1}{2}\%$  in molecular volume and about 1% in molecular refraction—are not the normal differences to be expected, but are bigger, because the equilibrium is different for that substance. They are larger differences than you would expect to find, for instance, in non-associated substances such as benzene and hexadiplobenzene.

The next point is that we should expect to find that the rates of reaction of the new hydrogen will be slower than the rates of reaction of the old, and to an extent which may easily lead to a variation in the ratio of 5 to 1 in the velocity constants of the reaction, and in exceptional types of reaction the differences may be very much greater. This is obviously of great importance, because there are many reactions, particularly in organic chemistry, where it is a matter of dispute whether a hydrogen atom takes part. If you can put one of the diplogen atoms in place of the hydrogen atom and measure the rate, that ought to tell you whether the hydrogen atom does take part in the reaction, or whether the reaction in which that hydrogen atom takes part is slow enough to affect the whole of the rate of the reaction.

Another point is that the discovery of this new isotope enables us to earmark the hydrogen atoms in a compound in very much the same way that Hevesy and Paneth have earmarked the atoms by the use of radioactive isotopes mingled with the other isotopes of the same elements which were not radioactive, but with this tremendous advantage, that whereas they were necessarily confined to the isotopes of the heaviest elements, which form part of only a very limited number of molecules, we are now able to do the same with hydrogen, which is a constituent of more chemical compounds than any other element.

All sorts of obvious applications arise out of this. One is in determining the readiness of reaction of hydrogen in ordinary groupings. Bonhoeffer has already shown that if you dissolve sugar in heavy water and let it crystallise out you get a redistribution, an almost equal partition of the heavy isotope

atoms between the water and the hydroxyl groups in the sugar, but that in those conditions you get no such partition with the hydrogen of the C—H groups of sugar. Sugar consists mainly of H O C H groups, one part redistributes with the water and the other does not redistribute at all. Again, Polanyi has found that DH gas reacts with water in the presence of a catalyst such as platinum black. It is clearly desirable to pursue this line of inquiry further, and it may be expected to throw much light on the nature of the processes involved. In particular, it will be important to find out how far the readiness of the hydrogen to undergo this change is affected in the first place by the nature of the atoms with which it is linked, and secondly—which is also a matter of great importance—by the nature of the other groups in the same molecule. That ought to throw a great deal of light on the nature of the chemical reaction.

To give another example of an important development, I may mention the question regarding electrolytic concentration. It is obvious to anyone who has seen the literature on the subject that when we know what the real difference of effective conditions is, the knowledge of that will throw great light on the intimate nature of these electrolytic processes themselves.

I have only tried to show a few of the directions in which we may expect the new reagent to be of value, but the most important services it will be found to render to science will be in directions which we do not yet suspect.

Dr F. W. ASTON, F.R.S. There are only two points which I wish to make. In the first place, I should like to emphasize how very small a discrepancy enabled the discovery of this element to be made. The measurement of the atomic weight of hydrogen by chemical and physical methods, as you know, compared with  $O = 16$ , is  $1.00777$ , and when I measured the relative masses of hydrogen 1 and oxygen 16 I got  $1.00778$ , and that measurement was made through helium, and therefore might have an accumulated error. The method I used was one in which one had to change the electric field. In the comparison of  $O^+$  with  $He^+$  I was able to use a small change of the field, as I could use the atom carbon as a marking point between the two, and I had only to change the field by 33%, but with  $He^+$  and  $H_2$  I had to change the field 100%, and, if there had been any systematic error, that on the two together might have been quite serious. I did estimate the possible error of the whole bracket as 1.5 parts in 10,000, and I thought it was considerably less.

There are three completely different measurements, the chemical atomic weight, the isotopic weight, and the abundance of the rare isotopes of oxygen

made by optical spectrographic observations. The accumulated error of those three experiments, of which in one the error might be admittedly 1.5 in 10,000, had to be less than 2 in 10,000 to prove that heavy hydrogen existed. I think Urey is to be congratulated as a brave experimenter to attack such a problem with such extraordinarily small differences in which to trust.

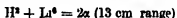
The only other point to which I desire to refer is that of nomenclature, which I think is extremely important. I was at the meeting in Chicago last June at which Urey and Lawrence were present. I was interested to note that the discussion threatened to become acrimonious, and I fear that our friends across the Atlantic may not like the suggestion coming from this side. I had a talk with Professor Bohr at the time, and he certainly at first insisted that the new isotope should be called hydrogen, he said it was not a new element, it had the atomic number 1, and therefore it was hydrogen. I admit that is true, but it is quite impossible to go on with the chemistry of this substance without having a new word for it, and at that meeting I suggested to Professor Urey that, although tradition allows the discoverer to name his discovery, it certainly does not guarantee that the public will use that name. We all know the story, for instance, of the "corpuscle," and in quite recent years there is the word "isotope," which is now so altered in meaning that its author would hardly know it. My own word, "mass-spectrograph," has also changed. I think we ought to have a word which will be as useful as possible, and therefore I strongly support Lord Rutherford's suggestion of "diplogen" and "diplon."

Dr P. HARTREE. First of all I should like to show you two experiments, illustrating the differences of vapour pressure from normal and heavy water and the different melting points of normal and heavy ice. According to Lewis and Macdonald, the vapour pressure of heavy water is smaller than that of normal water, at 25° C it is 12% less, and at 60° C about 8% less. Normal water has, at room temperature, a vapour pressure of about 24 mm., heavy water a vapour pressure of about 21 mm., a difference of 3 mm. At 60°, heavy water has a vapour pressure of 138 mm. against 150 mm. for normal water. I can show you this difference of vapour pressures with the help of a differential manometer. I have here a U-shaped tube containing mercury, on one side is a little heavy water and on the other side a little normal water, on the one side is the vapour of heavy water and on the other the vapour of normal water, and, at this room temperature, you will see a difference of 3 mm. in the levels. When the temperature increases, the difference will be greater. I have another apparatus of the same type which I plunge into hot water, and you can see as a result what a big difference there is!

The second experiment illustrates the difference in melting points. Normal water has a freezing point of  $0^{\circ}\text{C}$ , and heavy water, according to Lewis and Macdonald,  $3.8^{\circ}\text{C}$ . I have here two tubes, in one of which is a white pearl and in the other a blue pearl. They have been placed in a vessel where the temperature was about  $1^{\circ}\text{C}$ , and you will see that the heavy water is still in the form of ice, whereas the normal water has melted.

I should now like to make some remarks on the preparation of the heavy water.

Lewis and Macdonald\* have described the preparation of heavy hydrogen by means of the electrolysis of an alkaline solution with nickel electrodes. They do not give details of the experimental methods. There are, however, some precautions which must be observed if the enrichment of the heavy hydrogen ( $\text{H}^3$ ) is to proceed satisfactorily. This paper gives a description of a convenient method of working. For the working out of the method it was a big advantage that slightly enriched samples of water could be examined in the apparatus of Oliphant and Rutherford† in which lithium is bombarded with hydrogen ions. The long range  $\alpha$ -particles which are observed are due to collisions involving  $\text{H}^3$  according to the equation,



The amount of heavy hydrogen in ordinary water gives a measurable number of disintegration particles, and is a rapid method of estimating the amount of  $\text{H}^3$  at weak concentrations.

About 60 samples of enriched hydrogen, which had been prepared by electrolysis of normal water under varying experimental conditions, were tested in this way. When the method had been developed the  $\text{H}^3$  concentration was determined by density measurements.

The electrolytic cells were made of nickel supplied by the Mond Nickel Co. The cells were soldered carefully on the outside so that practically no solder came in contact with the electrolyte.

The electrolysis is begun with  $\frac{1}{2}\%$  NaOH. With a current of 150 to 200 amps and cooling water approximately  $10^{\circ}\text{C}$  the temperature of the electrolyte does not rise about  $25^{\circ}\text{C}$ . Grease and organic compounds in general must be carefully avoided. The wet surface is scrubbed with glass paper and steel wool and the electrolysis immediately begun.

\* 'J Chem Phys', vol 1, p 341 (1933)

† Oliphant and Rutherford, 'Proc Roy Soc', A, vol 141, p 359 (1933). Oliphant, Kinsey and Rutherford, 'Proc Roy Soc', A, vol 141, p 723 (1933)

More than 25 electrolyses have been carried out in this manner with four different cells without any case of failure. Since the initial concentration of alkali is so small it is possible to electrolyse to a  $\frac{1}{2}$  or  $\frac{1}{10}$  of the initial volume. The solution is then neutralized with  $\text{CO}_2$  and the water distilled off, as described by Lewis and Macdonald.

The efficiency of the electrolysis seems to be independent of the concentration of alkali over a wide range. No definite increase in efficiency was observed with increasing current density provided this was greater than 0.6 amp./cm.<sup>2</sup>. The temperature dependence of the electrolysis was not investigated since it was always necessary to work at a low temperature to keep down evaporation.

If the electrodes are really clean the efficiency has its full value from the beginning.

No.	$V_a$ cm. <sup>3</sup>	$V_b$ cm. <sup>3</sup>	$V_a/V_b$	% $\text{H}^3_a$	% $\text{H}^3_b$
1	3000	300	10	0.33	2.3
6	3000	100	30	0.33	5.3
15	40	1	40	12.2	100
23	2500	22.5	110	0.48	27
26	120	6	20	12	91.5

As soon as the concentration of  $\text{H}^3$  had the value of about 12% the mixture of oxygen and hydrogen coming off from the cell was burnt to water to avoid loss.

In the course of these experiments we have obtained enriched samples of heavy water containing from 30% to 98% of heavy hydrogen, corresponding in all to about 25 c.c. of pure heavy water.

Professor F. SODDY. The questionable extension of the now well-understood term "isotope" to heavy hydrogen and other supposed examples revealed only by band-spectra is, I think, probably responsible for the confusion, since 1927, between the chemical and physical (positive-ray) unit of atomic mass. The conception of chemically non-separable varieties of the same element, which dates from 1907-10, had its origin in the absolute chemical identity of radio-elements of different atomic mass, genesis, and modes of disintegration. It was followed by the displacement law of radioactive change, arrived at in its final form early in 1913, the critical evidence for which was based mainly on the work of Alexander Fleck in my laboratory at Glasgow. The double and single shift of place in opposite directions in the periodic table, which is produced by the expulsion of the  $\alpha$ - and the  $\beta$ -ray respectively from the disintegrating



atom, caused the word "isotope" to be coined. To me it was and still is a short term for the individuals of a group of two or more chemically identical elements existing together in constant natural proportion and separable only by the few physical methods that depend directly on the masses and therefore the velocities of the molecules.

In the country of its origin at least this independent experimental conception of chemistry has always been overshadowed by the contemporaneous conception of the nuclear atom of Rutherford and Bohr, and the conception of atomic number of van der Broek, as later determined experimentally from X-ray spectra by Moseley, ultimately for all the elements except those of the first short period. The positive ray method of Thomson, Aston's extension of the conception of isotopes to the non-radioactive elements, and his discovery of the whole number rule, furnish now the best proof of the correctness of the conception itself, as it was arrived at in the narrower field of radio-chemistry, and particularly the constancy of the natural ratio. The only slight exception with regard to the second criterion, the constancy of natural ratio of isotopes, is boron, examined by Professor Briscoe and others, which is, as Aston has pointed out, most favourable for separability by physical processes such as diffusion.

Up to 1929, the chemical and physical methods of determining atomic weights led to the most gratifying agreement for the four elements up to and including oxygen, found by Aston by the mass-spectrograph to be "pure," and for which some of the finest atomic weight data are available. Thus, in terms of oxygen 16, the chemical and physical (mass-spectra) values were as follows —

Nitrogen	14.008 and 14.008
Carbon	12.0025 and 12.0036
Hydrogen	1.00777 and 1.00778

Then, in 1929, from evidence of band spectra, the existence of small proportions of the two isotopes  $O^{16}$  and  $O^{17}$  was inferred. If these are really isotopes in the original sense, their existence demands the existence also of very nicely equivalent proportions of corresponding isotopes in the other three elements, and sure enough, when looked for, band spectra indicated also the existence of  $N^{14}$ ,  $C^{12}$ , and  $H^1$ .

The first point is, and it has not been mentioned to-day, that the heavy hydrogen does not appear to be in nearly sufficient proportion to bring back the concordance between the chemical and physical numbers. I think Aston

refers to 35,000 to one, and Lord Rutherford to-day gave it about half that.

Lord RUTHERFORD I said 6500

Professor SODDY 6000, at any rate, is required to bring it back to the right value, and how these proportions can possibly be determined seems to me very difficult to believe, I mean, for these isotopes at any rate it is mere guesswork, and I think the knowledge of what is required to bring the value right must have a certain weight—unconsciously, no doubt—with the investigator. Further, it has been forgotten that this marked difference of chemical character that has been discovered for heavy hydrogen entirely destroys the basis of the prediction. To any chemist who is acquainted with the most extraordinarily careful atomic weight work on which the ratio of hydrogen to oxygen was founded, it is incredible that a substance so easily separable can have had any more effect on the chemical than it does on the physical ratio. One need not go into the question of whether heavy hydrogen is or is not absorbed by palladium or the usual methods by which the chemists purified their gases. Either hydrogen and heavy hydrogen were separated completely or not at all. Otherwise the differences in purification must inevitably lead to differences in the atomic weight determinations. The fact that this value of the chemists was a concordant value, although the two substances are readily separable, seems to me conclusive evidence against the validity of the original prediction. To me it is a comedy of errors, it is not a question of prediction at all.

With regard to separability, the instance would appear to be much more analogous to the separation of a pair of homologues, such as zirconium and hafnium, rather than isotopes. As to the criterion of constant ratio, I do not know what the evidence is, but I have yet seen none which would exclude the possibility that we are dealing with substances analogous to oxygen and ozone, that is, that heavy hydrogen is not in natural constant proportion in hydrogen at all, but is formed by the electric discharge or by electrolysis of alkaline solutions under high pressure and current density. Aston quotes in his recent book, "Mass Spectra and Isotopes," the evidence of Sandford, that the band due to  $\text{C}^{12}\text{C}^{13}$  is abnormally intense in the spectra of certain types of stars, and for what stellar evidence is worth to the chemist that shows that these are not isotopes in the original sense of the word.

The relative ease with which the light elements are transmuted apparently ought also to make for caution in applying properties inferred from the behaviour

of the heavier elements to those of the first short period, and particularly to hydrogen. I have never given a place in the periodic table to this element at all, and therefore regard the expression "hydrogen isotope" as doubly a misnomer. It may prove that it is even easier to synthesize heavy hydrogen artificially than to disrupt the light elements.

I do not wish to detract in the very least from the intense interest and importance of this discovery. I think it is of fundamental importance, and the Americans are to be congratulated on it. I recall that two Americans, McCoy and Ross, in 1907, when confronted with a pair of isotopes, had the courage to call a spade a spade and to declare that radiothorium and thorium are chemically inseparable by any process. Heavy hydrogen bids fair, I think, to be one of the great discoveries of the century. I hope it may prove valuable not only in smashing up a few atoms, but even more so in smashing some of the physical theories of the present day about the whole of them!

Dr. M. POLANYI: I should like to say a few words about the experiments of Dr. Horvut with hydrogen containing a small percentage of diplogen which have been made with a view to measuring the rate of ionization of diplogen, in connexion with the problem of over-voltage and catalysis. If hydrogen containing 1% of diplogen is put into a 100-c.c. container and 10 c.c. of water is added, and we add half a gramme of platinum black and shake up the whole at ordinary temperature for 10 minutes, the 1% of diplogen is reduced to  $\frac{1}{4}$ %. This proves the existence of a catalysed interchange between the diplogen and the water.

It is well known that platinum is a good hydrogen electrode, and that the function of the hydrogen electrode consists in bringing about equilibrium between hydrogen and the hydrogen ions of the solution in which it is immersed. In doing so, the current is caused to pass through the electrode carrying hydrogen atoms in the form of hydrogen ions into the solution, and hydrogen ions at the same time and in the same quantity pass from the solution and form hydrogen. If part of the hydrogen on the electrode is substituted by diplogen then these two currents tend to interchange the hydrogen and the diplogen. On half a gramme of platinum black there is about 0.2 ampere of current going to and fro in both directions.

We have carried out a few experiments to make sure that what we are measuring here is really the actual rate of ionization and not some accidental factor such as diffusion. We changed the solution, taking an acid solution, then an alkaline solution, and then a different acid as well as alcoholic solutions,

and we found that as we did so the rate of ionization changed quite considerably, in particular the rate of ionization was very much slower in the alkaline solution than in pure water. If we take the rate of ionization in water as 1,  $n/1$  hydrochloric acid is 0.7,  $n/1$  sulphuric acid 0.2, and  $n/4$  NaOH 0.4, in alcohol with 2% water the rate is 0.4 and if we add some alkali to this solution, the ionization disappears, so that we could not measure it in the time used for these experiments.

This shows that we actually measure not the diffusion, which, of course, could not be affected in this way, but the ionization process itself. In addition, I think it throws some light on the question of what is the inertia attached to the ionization of hydrogen. It is well known that, when hydrogen is formed on an electrode, it is necessary to have a certain over-voltage to force the hydrogen out of the solution, and from this over-voltage one can conclude that there is some chemical or physical resistance which has to be overcome. Evidently, this resistance is identical with the inertia attached to the ionization of hydrogen, which we have measured in our experiments. We recognize that it depends on the nature of the solution in which the electrode is immersed. This indicates very strongly that of the two theories which have been current about the nature of this resistance, at least for platinum, only one can hold good. One of these theories assumes that it is the dissociation of the hydrogen into atoms, which is accompanied by inertia, and that this caused over-voltage. The other theory assumes that the hydrogen instantaneously decomposes into hydrogen atoms, and that it is the second step, the ionization of the hydrogen from the atomic state to the ionic state, to which over-voltage is due. It is evident that the first step, which is really a reaction between hydrogen and platinum, could scarcely be influenced by the nature of the solution, whereas the second step would naturally be strongly influenced by it. Therefore, at least, for platinum, the cause of over-voltage must lie in an inertia attached to the transition of the hydrogen ion from the solution to the atomic state in its adsorbed form on the platinum.

It seems to be generally assumed that deuterium will always react more slowly than hydrogen. As I may partly be responsible for this view,\* I should like to point out that this is not always correct. Lower reactivity of deuterium compared with hydrogen results mainly from two causes: (1) the existence of zero point energy, and (2) the quantum mechanical leakage of particles through energy barriers. Whilst the leakage through the barrier is always

\* Cremer and Polanyi, 'Z. phys. Chem.,' B, vol. 19, p. 443 (1932). See also Eyring 'Proc. Nat. Acad. Sci. Wash.,' vol. 19, p. 78 (1933).

greater for the hydrogen than for the diplogen atoms, the effect of the zero point energy may occasionally favour the reverse ratio. I will confine myself to one special case, as the general treatment will be published shortly by Bawn and Ogden. Compare the reaction of a free hydrogen and a diplogen atom, in the initial state the atoms possess no zero point energy and their energies will be equal. However, at the top of the barrier there will be a zero point energy present,\* and this will be greater for the complex reacting with the hydrogen atom than for that reacting with the diplogen atom. The effect of the zero point energy at the top of the barrier is, therefore, to increase the activation energy of the hydrogen atoms to a greater extent than that of the diplogen atoms.

Professor E. K. RIDEAL, F.R.S. I cannot do more, in the time available, than give you a very brief résumé of the work which has been carried out by Drs L. and A. Farkas during the last few months. They had been occupied, when in Germany, in Professor Haber's laboratory in Berlin and Dr. Bonhoeffer's laboratory in Frankfurt, with experiments on the ortho-para hydrogen conversion, and it did seem possible that the thermal conductivity methods used for studying the ortho-para hydrogen system might be applicable to the diplogen-hydrogen system. The only method we have had of estimating the quantity of diplogen in hydrogen has been that of converting it into water and determining the density, a laborious process and one requiring at least several milligrams, whereas with the thermal conductivity method 0.002 milligram is all that is necessary.

The experimental arrangement was a slight modification of that described in a recent publication by Dr. A. Farkas for the sensitive measurement of ortho-para hydrogen composition. This method is based on the different slopes of the specific heat curves for the two hydrogen isotopes.  $H_2$  has at 80° K. a specific heat of 3 cal./mol. which rises to 5 cal./mol. at room temperatures, whereas the specific heat of  $D_2$  (and also that of HD) does not alter much over this temperature range and is about 5 cal./mol. The actual determination of the hydrogen diplogen concentration was carried out by measuring the temperature of an electrically heated wire stretched in the hydrogen at about 0.04 mm. pressure. The difference in the temperatures of the wire which were reached when pure  $H_2$  and pure  $D_2$  were placed in the conductivity cell amounted to about 20° C. corresponding to a change of 6 ohms in the resistance of the wire. The calibration was effected by the samples of water of different

\* Eyring and Polanyi, 'Z. phys. Chem.', B, vol. 12, p. 279 (1931)

D content We have to thank Dr Harteck for having performed the very tedious process of concentrating to its last stages our heavy water

The next point was to study the equilibrium  $H_2 + D_2 \rightleftharpoons 2HD$  If the hydrogen and diplogen mixture is heated on a catalyst, in fact on a nickel wire, that equilibrium is rapidly established The formation of HD from  $H_2$  and  $D_2$  effected a rise of approximately 1 ohm in the resistance value of a 50% mixture, of  $H_2$  and  $D_2$  and this 1 ohm corresponds to the establishment of an equilibrium when 50% HD is formed and 25%  $H_2$  and 25%  $D_2$  disappeared

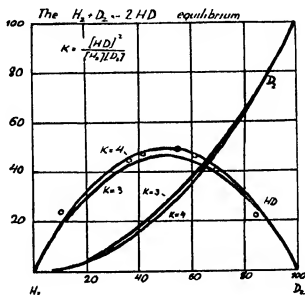


FIG 1

By making suitable mixtures of hydrogen and diplogen and bringing them to equilibrium, one can find the equilibrium constant In fig 1 are the experimental values, and one can see that the equilibrium constant lies between 3 and 4, the curves correspond to the theoretical values The first point that has been established here is that, in accordance with theory, the equilibrium constant does not change much above room temperature and up to 600°, and, secondly, that the equilibrium constant is very close to 4, as was also to be anticipated by theory

The next point to consider is how this equilibrium is established We have seen that on catalysts it can be established very readily at quite low temperatures How is it established in the homogeneous gas phase at high tempera-

ture? It was found that there was no reaction at all in quartz under 600° C. At higher temperatures, however, a reaction took place, and samples could be removed and analysed and we can get the order of the reaction.

Fig 2 shows the reciprocal time of half change as a function of the pressure at 725° C. The order of the reaction given by this curve lies between 3/2 and 2.

There are two possible mechanisms for this reaction. We can first of all imagine a molecular reaction  $H_2 + D_2 \rightleftharpoons 2HD$ , and secondly an atomic reaction giving  $H + D_2 = HD + D$ ,  $D + H_2 \rightleftharpoons HD + H$ , where H and D are formed by the thermal dissociation of  $H_2$  and  $D_2$ . The velocity of the molecular reaction will be proportional to the square of the pressure ( $p^2$ ) and of the other proportional to  $p^{3/2}$ . From the velocity of the reaction at different

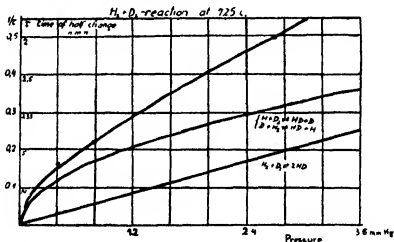


FIG 2

temperatures, it is possible to split up the curve into two corresponding to each of the above reactions and to get the variation of the collision efficiency with the temperature, and from that what is called the energy of activation. It is found that for the  $p^{3/2}$  reaction, the one going through the atoms, the energy of activation is 50,000 + 7,000 calories, i.e., 57,000 calories, where 50,000 calories are used up for the thermal formation of the H and D atoms. That is in good agreement with the result obtained for the thermal parahydrogen reaction which proceeds according to the reaction  $H + H_2 \rightarrow H_2 + H$ . The molecular interchange reaction has an energy of activation of the same order, that is, between 50,000 and 60,000 calories.

Drs Farkas have also examined the reaction with water

Table I —  $D_2 + H_2O$ -reaction. 1 mm.  $D_2 + 17$  mm.  $H_2O$ 

Temperature	D-content after 2 hours reaction.	Collision efficiency
°C	%	
500	98	2 $10^{-14}$
600	90	2 $10^{-12}$
675	76	9 $10^{-12}$

From the data of Table I one can show that the energy of activation of the reaction is again of the order of 60,000 calories

One can analyse the gas mixture from time to time and find out how these particular reactions are proceeding. We have two possible reactions,  $H_2O + D_2 = D_2O + H_2$ , and  $H_2O + D_2 = DHO + HD$  and it is found that the ratios of the velocities of these two reactions are of the order of 1 : 3

It is also interesting to note that the collision efficiency, extrapolated to room temperature, is about  $10^{-12}$ , which indicates very clearly that the so-called Oliphant reaction does not proceed directly, but proceeds probably through a catalyst, as shown by Dr. Polanyi.

Some rather interesting things were discovered in the examination of the effusion of hydrogen deuterium mixtures through a fine nozzle, as the effusion may cause a very great change in the composition of the gas.

In Table II are shown the changes in the composition of the residual gas when the pressure  $p_0$  (several hundredths of a millimetre) is diminished to a pressure  $p$ .

Table II

$p_0/p$	% $D_2$ observed	% $D_2$ calculated
1	47	—
1.5	50.7	50.7
2.0	53.0	53.0
3.25	56.0	57.8

The observed results are in very good agreement with the values calculated on the basis of the theoretical equation

$$\left(\frac{H_2}{D_2}\right) = \left(\frac{p_0}{p}\right)^{s-1}$$



where " $s$ " the separation factor, is the ratio of the molecular velocities,  $\propto \sqrt{2}$

The diffusion of the two isotopes in their passage through palladium, which is one of the best methods of purifying hydrogen, was also studied

If one starts with hydrogen and deuterium mixtures the equilibrium between all three is readily established. The rate of diffusion is temperature exponential, and proportional to the expression  $e^{\frac{18000}{RT}}$ , where 18,000 cal is the activation energy for the diffusion. If we analyse the hydrogen deuterium of the diffused material and compare it to the hydrogen deuterium ratio of the original material  $\frac{(H/D)_{\text{diff}}}{(H/D)_{\text{orig}}}$  we find a very interesting thing. This ratio at low temperatures

is relatively high,  $\propto e$ , 1.5 at 160° C, while as the temperature rises the ratio gets closer and closer to unity, in other words, the deuterium, as it were, remains behind, and the light hydrogen passes through more freely. The expression for the ratio as a function of the temperature is given by  $e^{\frac{830}{RT}}$  where 830 calories, represents the difference in activation energy for some process in the diffusion. It is remarkable that the observed 830 calories is very close to the energy difference in the zero point energies for the two substances.

This observation suggested that one should be able to separate hydrogen and deuterium in those chemical reactions in which the velocity whatever the energy of activation happened to be, is sensitive to changes in the energy of activation of  $\approx 1000$  calories. For example, if one takes metals and dissolves them in water or dilute acids one finds the energy of activation for the process is extremely small. One would therefore anticipate that one could obtain, coming off in the gas, a mixture of hydrogen and deuterium, but that the ratio of hydrogen to the deuterium should be much higher than in the water itself. That in fact occurs, and I have here a few tentative figures for the analysis of gas coming off from a 30% hydrogen-deuterium water. If one takes zinc, for example, with a slight quantity of sulphuric acid, and dissolves that in this water, one finds that the ratio of hydrogen to deuterium in the effluent gas is 3.5 to 1, in other words, one has almost the separation that is found by electrolysis. Other metals are not so effective. For calcium it is 1.5 to 1, for sodium 1.2 to 1, and for aluminium 2 to 1. There is a number of other possible methods of effecting this separation by chemical means, I have simply shown this one because it is very simple, and possibly lends itself to technical application.

As I said before, this is just a very brief summary of the work that the two Drs. Farkas have been doing since they came from Germany.

Professor R H FOWLER, FRS Personally, I have been interested in trying to understand how it is that electrolysis separates the hydrogen and the deuterium in such a way that over very wide ranges of hydrogen-deuterium concentration in the water that you are electrolysing you get a rate of evolution of hydrogen and deuterium which is some factor,  $Q$ , times the ratio of their concentrations in the water. I have been trying to satisfy myself, for example, by general arguments that the ideas put forward by Dr Polanyi which are probably right, represent the only possible way of talking about that separation. In doing that, I tried to avoid as far as possible all reference to details of the mechanism, and merely to see what was required to get such a constant evolution rate independent of the concentration ratio in the solution, at the same time recognizing that you have a *steady* process of electrolysis going on, that you have no accumulations of hydrogen or other substance anywhere except in the evolved gas at the expense of the solution in general.

The point that I wish to make is this. If you consider the whole electrode system where the process is going on, hydrogen is being fed into that by various processes, by the current carrying process, for example, from the solution in general, and is being evolved in gas bubbles, and those two things must take place at the same rate. From a simple argument such as that you can, I think, get a certain control over your theories which it is well worth while to develop at some length. I do not, propose to do that now, I shall merely indicate the sort of arguments employed.

For example if diffusion in your liquid were negligible and your only hydrogen carrying process was the actual current transported by the hydrogen ions then, no matter what your electrode processes might be, the ratio of the rates of evolution of the hydrogen and deuterium would be just  $Q$  times the ratio of the concentrations, where  $Q$  is simply the ratio of the mobilities—that and nothing else. Now, that is not true, and what is happening is that atomic diffusion in the neighbourhood of the electrode prevents the building up which would be required by that result, and actually the boundary there is kept by diffusion at a fair sample in H/D ratio of the whole liquid in bulk, in spite of the transport of H and D to the electrode.

If you go further into the process and consider the exchange of hydrogen between the electrode itself and the boundary layer of the liquid, you have a deposition of hydrogen on the electrode from the boundary layer, this deposition is a current-carrying process, controlled by the over-potentials, which may be different for the H and D, and that by itself could give rise to a ratio of rates of evolution of the two gases of the observed form, provided this

deposition is the only process effective in exchanging hydrogen and diplogen between the solution and the electrode. The factor  $Q$  is then just the ratio of the rates of deposition at the given concentration and under the given conditions of the electrolysis. That would be true no matter how the atoms after deposition reacted to become gas molecules, but, as Dr Polanyi pointed out in his interesting experiments, that is not the only process of exchange. The current-carrying process is not the only process of exchange of hydrogen between the liquid and the electrode, and other conditions come in which must be taken into account. When you take those other conditions into account, of course, it may very well happen, and will happen at low current densities, that the current carrying process is trivial, and you still have your equilibrium set up between the adsorbed hydrogen and the hydrogen in the water, and then it matters very much what happens later on. The process which controls the evolution will then depend on the rate at which the adsorbed hydrogen react atoms in pairs to form the evolved gas.

When you trace that through you find that a certain condition has to be satisfied between the efficiency of interaction of HD, HH and DD, but a relationship which is likely to be true, and, if that is true, you then still get a ratio of rates of evolution of the form  $QH/D$  over an infinite concentration range, but your factor  $Q$  may depend here not merely on the overpotential factors but also on a collision-efficiency factor between the adsorbed hydrogens. The question therefore seems to me still open as to what part of the observed  $Q$  comes from one or other of those processes. This is not the place to go into such a special point more deeply, I merely wish to point out that such questions do arise and will need further interesting discussion as the facts become better known. Such discussion will inevitably lead to a better understanding of the whole phenomenon.

Mr R P BELL. I have a very short report to make on some experiments carried out by Mr Wolfenden and myself on the effect of different factors on the electrolytic fractionation of water. We determined the concentration of diplogen by measurements of specific gravity, independent determinations by different observers agreeing to within 1 part in 100,000. Water of known diplogen content was electrolysed under different conditions to a known fraction of its original volume and the concentration of diplogen in the final product determined by a specific gravity determination. We have expressed the efficiency of separation by the factor  $\alpha$  in the equation

$$d \log D = \alpha d \log H,$$

i.e., our  $\alpha$  is the same as Professor Fowler's  $Q$ .

Our first experiments deal with the effect of varying the cathode metal, and the results are given in Table I

Table I—Effect of Cathode Metal

D/H = 0 1% → 0 3%

Electrode	Electrolyte	$\alpha$
Ni	1% NaOH	0 22
Ni	8% NaOH	0 20
Pt	1% NaOH	0 19
Cu	1% NaOH	0 10
		} $\pm 0 05$

The estimated possible error in the value of  $\alpha$  is 0 05, this, however, refers to the absolute values, and the differences between the values in a series should be correct to about 0 02. Within the experimental error, exactly the same efficiency was obtained using nickel, platinum and copper as cathodes. In one experiment the initial caustic soda concentration was 8%, and hence the final concentration was something like 40%. It has been reported that the separation becomes much less efficient in concentrated alkali. This is not confirmed by our results, the efficiency being just the same as in dilute solutions.

Table II shows the result of experiments on the effect of temperature and of current density

Table II—Effect of Temperature and Current Density

I D/H = 0 25% → 0 5%

Electrolyte	Temperature	$\alpha$
2% NaOH	100°	0 27
2% NaOH	10°	0 24
		} $\pm 0 05$

II D/H = 0 05% → 0 15%

Electrolyte	Current density	$\alpha$
1% NaOH	10 amp./cm <sup>2</sup>	0 18
1% NaOH	0 08 amp./cm <sup>2</sup>	0 27
		} $\pm 0 05$

It will be seen that there is practically no difference between the efficiencies at 10° and 100°. There does appear to be a slight difference in  $\alpha$  in the two current

density experiments, but it is doubtful whether it is greater than the experimental error. These experiments need repeating with greater accuracy.

These are the experimental results which I wish to report, and I think that the most interesting point is that the separation is much less sensitive to different variables than one would have expected. It has been suggested that the separation is in some way connected with over-voltage phenomena. If this is so, one would expect some change in efficiency on changing the nature of the cathode metal, actually nickel, platinum and copper were found to be equally efficient. The absence of any temperature effect is also unexpected, though it may be that the processes taking place in the neighbourhood of the cathode are not much affected by the temperature of the bulk of the electrolyte.

Our average value for the efficiency  $\alpha$  was about 0.2, agreeing with the value found by Lewis. Dr. Harteck's results also appear to indicate about the same efficiency.

Mr J. D. BERNAL. Through the kindness of Lord Rutherford, I was able to get some of the 91% heavy water. This was put into a small capillary, and a single crystal of ice was made and photographed\*. The resulting cell measurements showed a slight, but I think, genuine difference from the values for ordinary ice taken under similar circumstances and with the same camera. The  $a$  axis was 4.50 Å for heavy ice as compared with 4.52 Å for ordinary ice, and the  $c$  axes 7.36 and 7.39 Å respectively. The difference is small, and amounts in specific volume to a little over 1%, but it is very interesting to note that, if this result is correct, the volume difference is in the opposite sense to the volume difference for water. The apparent molecular volume of heavy water is slightly greater than that of ordinary water, the apparent molecular volume of heavy ice is slightly smaller than that of ordinary ice.

I think I can give a theoretical reason for that being so, and at the same time point out that there are two quite different reasons why diplogen compounds should differ from ordinary compounds. One has already been mentioned: the difference between the zero point energies. The other is the difference which is a pure mass effect, due to the double mass of the diplogen in the water, and that affects the whole nature of the water in this quite simple way. The nature of the difference of the so-called polymers of the water is really determined by the orientation, by the regular or irregular arrangement of the molecules.†. The moment of inertia of the heavy water is about twice that of

\* The photographs were taken and analysed by Miss H. Megaw.

† Bernal and Fowler, 'J. Chem. Phys.' vol. 1, p. 55 (1933).

ordinary water, and therefore it is in general a slower rotator, so that at the same temperature heavy water should be more ice-like, that is, its co-ordination number should be lower, giving it a larger volume. That is what you get in the actual waters, you do get a larger volume for the heavy water. On the other hand, with ice, where the rotation is completely suppressed and all the atoms are in regular positions—suppressed, that is, but for a small number of rotators to account for the large dielectric constant—you get a smaller volume.

All the properties of heavy water fit in with this hypothesis. The most interesting is the relation of surface tension and viscosity. The viscosity of heavy water is definitely 30% to 50% larger than that of ordinary water, whereas the surface tension of heavy water is actually only 0.8 of the surface tension of ordinary water. Here you have two phenomena both associated with polymerization and going in different directions. The abnormally large viscosity of heavy water is not a translatory but a rotary viscosity. The ease with which an object can be moved through water depends on the degree of spontaneous rotation of the molecules. That will be small for heavy water and therefore you have higher viscosity. On the other hand, the surface tension of water depends on the force between the molecules and is not directly affected by the rotation and therefore you would expect it to be the same or somewhat smaller for heavy water. It is possible to construct a theory for these effects, but I cannot go into it here.

The differences due to this mass effect will be very apparent in certain compounds where the other effect may not be so noticeable, and particularly where the possibility of the mobility of the hydrogen ion is concerned, especially in acids. I think the general theory of the mobility of the hydrogen ion depends, at any rate in the form of which Professor R. H. Fowler and I have worked it out, very much on the mass, and the difference in mobility shown by the hydrogen ion is about 50% greater than could be accounted for by a normal viscosity effect.

Dr W. JEVONS. Spectroscopic investigations in connection with the hydrogen isotope have not been reviewed in this discussion, beyond Lord Rutherford's opening remarks on Urey, Brickwedde and Murphy's original observation and measurement of the weak Balmer lines of D (i.e.,  $H^2$ ) which accompanied the much stronger corresponding lines of H (i.e.,  $H^1$ ), and Hertz's conspicuous success in obtaining D in such a high state of purity that the line spectrum showed no trace of the Balmer lines of H and the molecular spectrum

consisted of band-lines of  $D_2$  with no trace of those of  $H_2$  and  $DH$ . I therefore venture to draw attention to the significance of the spectroscopic results so far secured

These measurements of the Balmer lines of  $D$ , and Ballard and White's subsequent measurement of six lines of the Lyman ultra-violet series of  $D$ , constitute the only examples of an observed electronic isotopic effect which is given directly by Bohr's well-known expression for the Rydberg constant for one-electron atoms,  $H$ ,  $He^+$ ,  $Li^{+1}$ ,  $Be^{++}$ , (For atoms with two or three electrons, such as  $Li^+$  and  $Li$ , the observed isotope effects are, of course, less ample in theory, the theory has been given by Hughes and Eckart)

Turning now to band-spectra of diatomic molecules we have so far records of the  $H$  isotope effect in three well-known band systems of  $HCl$ ,  $H_2$  and  $OH$  --

(i) It is now well known that in the infra-red  $HCl$  absorption band near  $\lambda 3.46 \mu$  each strong line of the more abundant molecule  $HCl^{35}$  has, on the low-frequency side of it, a weaker line of the less abundant molecule  $HCl^{37}$ , that is to say there are two overlapping bands arising from the  $Cl$  isotope effect. Hardy, Barker and Dennison have observed near  $\lambda 4.8 \mu$  the corresponding pair of bands due to  $DCl$ , with a similar duplex structure arising from the  $Cl$  isotope effect, no less than 19 lines of the molecule  $DCl^{35}$  and 17 lines of the rarer molecule  $DCl^{37}$  being measured. The vibrational displacement is very large ( $DCl - HCl$ ,  $\delta\nu = -805 \text{ cm}^{-1}$ ), the coefficient  $\rho - 1$  being very much greater than for any other observed isotope effect. The deduced mass ratio is  $H : D = 1.00778 : 2.0137$ , in close agreement with the mass spectrograph determination made earlier by Bainbridge, namely,  $1.00778 : 2.01363$  (on the chemical scale  $O = 16$ ). The estimated abundance ratio  $H : D = 35000 : 1$ , has been superseded by Bleakney and Gould's result,  $5000 : 1$ . The ratio  $6000 : 1$  quoted by Lord Rutherford is presumably a still later and closer estimate.

(ii) Using a gas mixture consisting of about 25% of  $H_2$ , 50% of  $DH$  and 25% of  $D_2$ , Miss M. Ashley and Professor G. N. Lewis have obtained 21-foot grating spectrograms, and identified and measured several hundreds of new lines. Many of these lines have been arranged into P, Q and R branches of  $DH$  and  $D_2$  bands which correspond to the  $\alpha$  bands ( $3p^2\Pi \rightarrow 2s^2\Sigma$ ) of  $H_2$ . The large isotopic displacements observed are in satisfactory agreement with those calculated from  $H_2$  data, and include, not only the usual vibrational and rotational displacements, but also very considerable electronic displacements, the latter amounting to as much as  $2.4 \text{ cm}^{-1}$  for  $DH - H_2$ . Alternating line intensities are observed in the bands of the homonuclear molecule  $D_2$ ,

but not, of course, in those of the heteronuclear molecule DH. The alternations in the  $D_2$  bands are such that, without reasonable doubt, the spin quantum number for the D nucleus (dipion) is 1, which is twice that of the H nucleus (proton).

(iii) In the 0, 0 band of OH,  $^2\Sigma \rightarrow ^2\Pi$ , at  $\lambda 3084$  the hydrogen isotope effect has been reported, though somewhat tentatively, by Miss K. Chamberlain and H. B. Cutter, who observed and measured several new lines in the positions computed for lines of the  $^2R_{21}$  branch of the corresponding band of the isotopic molecule OD. This result is confirmed and greatly extended in a detailed investigation by Johnston and Dawson, who have identified and analysed large portions of the 0,1, 1,0 and 2,0 bands as well as the above branch of the 0,0 band. The observed displacements include pronounced isotope effects on the spin-doubling in the  $^2\Pi$  state in addition to the larger vibrational and rotational displacements. It may reasonably be expected that this investigation will lead to a very exact value of the H/D mass ratio, and also reveal yet unobserved isotope effects on the A-type doubling in the  $^2\Pi$  state and the spin-doubling in the  $^2\Sigma$  state.

LORD RUTHERFORD, O.M., F.R.S. I am sure you will all agree that we have had a most interesting meeting, and I regret that lack of time does not allow many speakers to contribute to our discussion. At the moment, only a few scientific men in the country have even a cubic centimetre of heavy water for experimental purposes. I hope this difficulty will soon be removed and that our industrial friends will help us to obtain supplies of the new water at a reasonable price. The cost of making the new water in places like Cambridge where the cost of power is high, is almost prohibitive. If a further discussion is held again, say, in six months' time, new information is likely to be available on the biological as well as on the chemical/physical side.

Before closing the discussion, I am sure you would like me to say a word on the points raised by Professor Soddy in his communication. As you all know, Professor Soddy was the discoverer of isotopes in the radioactive bodies and coined this name because the radioactive elements appeared to occupy the same place in the periodic table, and to be inseparable by chemical methods. Much water has flowed under the bridge since he made that discovery and we now speak of the isotopes of an element not as inseparable bodies but as consisting of atoms which have the same nuclear charge but different masses. I think no one will doubt that heavy hydrogen has the same nuclear charge as hydrogen with one electron circulating round the nucleus, and from this point



of view is properly to be regarded as an isotope of hydrogen. I, of course, quite understand the reason of Professor Soddy's difficulties. The name isotope was first applied to atoms of the heavy radioactive elements differing only slightly in mass, chemically indistinguishable and very difficult to separate even by diffusion. The name is now given to the atoms of heavy hydrogen which can be readily separated from ordinary hydrogen and shows some very distinctive properties. This contrast is due to the fact that in hydrogen the masses of the isotopes vary in the ratio of 1 to 2, and are thus readily separated by diffusion or by processes like electrolysis where the mass plays a large part. Even the radioactive isotopes could be partially separated by diffusion methods but owing to the small difference in their masses the process would be a slow and tedious one.

I hope that I have been able to convince Professor Soddy that in using the word isotope for heavy hydrogen we are not contradicting the essential ideas involved in his first use of the word.

### *The Kelvin-Poincaré Problem of Stellar Evolution*

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### *On Dynamical Evolution*

By SIR JOSEPH LARMOR, F.R.S.

#### *Summary*

The evolution of a rotating liquid mass with an assigned angular momentum about a central axis has been sketched by Poincaré in his celebrated memoir on a rotating liquid mass (*Acta Math.*, 1885). The liquid continually contracts so as not to violate the principle of degradation of mechanical energy which we take for the criterion of the trend of evolution. For the sake of mathematical simplicity we make the assumption that the contraction permits the liquid to be always homogeneous. Initially a sphere with a "small" angular velocity about a diameter, the liquid mass acquires greater and greater velocity as it contracts with constant momentum and passes through the stable

Maclaurin and Jacobian forms. This is the order up to the critical Jacobian ellipsoid, if for conserved momentum the total mechanical energy becomes continually smaller so that the principle of degradation of energy is not contradicted. The next type in the succession, a pear-shaped figure, is unstable, and so it is presumed that a cataclysm involving tumultuous change sets in at this stage. In view of the slowness of the evolution it is perhaps to be expected that the cataclysm would be smooth in its earlier stages. We examine the general features of these earlier stages and show that contraction subject to degradation of energy permits the Darwin-Poincaré pyriform solution, unlike that of Jeans. The former seems to have been found to be stable while the latter, the more developed pear, has been shown by Jeans to be unstable. Without discussing the cataclysm further, we pass next to a series of double-figure configurations (both components spherical), which is characterized by two parameters,  $\lambda$  the mass-ratio and  $r$  the separating distance. In the diagram of the course of evolution which summarizes the investigation all the configurations are brought in. If the cataclysm ends in such a smooth state of fission, we must have  $\lambda < 3.4$  which is in keeping with the observation of double stars. As regards the earlier stages, the Maclaurin spheroid shrinks in every direction, but not the Jacobian ellipsoid.

# 1 Introduction

The historically famous problem of gravitating liquid masses is discussed here from a point of view which has not received due attention, since Lord Kelvin published a brief statement of results, without proof\*. As discussed here the problem is this: given a mass  $M$  of a homogeneous liquid rotating uniformly like a rigid body, with a fixed angular momentum  $\mu$ , what will be its course of evolution, if it slowly contracts so as always to remain homogeneous? Owing to the recent progress of astrophysics the interest in this form of the problem is now mainly historical. Nevertheless, there is a good reason for presenting these considerations, as Sir George Darwin, who had this problem in view, had to leave it undiscussed, for the question of the stability of the pear-shaped figure still needed attention when he died, and when the last word was said in this connection, the interest in the problem of evolution had evaporated.

There is also another reason, equally good. The current idea about the origin of the binaries is essentially that held by Darwin and Poincaré†. But

\* "Thomson and Tait," 2nd ed. § 778, vol. 2 (1883)

† "Acta Math." p. 379 (1885) see also Jeans's "Astronomy and Cosmogony," p. 414 (1929)

there does not seem to be on record a model of evolution, in conformity with the old or the new ideas of astrophysics, which would support this current view, or otherwise. It may therefore be worth while developing fully the Kelvin-Poincaré stellar model.

### 2 *An Account of several Successions of Figures of Equilibrium.*

Let us take a rectangular set of axes  $Oxyz$  fixed in space.  $O$  is the centre of a homogeneous liquid sphere of mass  $M$ , radius  $a_0$  and density  $\rho_0$ . Let a spin  $\omega_0$  be given about the axis  $OZ$ , being small enough for a spheroidal form to be possible. We will suppose that the mass is contracting, owing to cooling, and that the contraction is so slow that whatever form the mass assumes it is always homogeneous. As the system remains isolated, the angular momentum  $\mu$  as well as the mass  $M$  must remain constant. Throughout this paper the liquid will be supposed to be homogeneous. We will consider a succession of figures of equilibrium all of which have the same mass  $M$  and angular momentum  $\mu$ . When tables are constructed for the densities, angular velocities, etc., corresponding to these figures, it should be possible to derive a sequence of configurations characterizing the course of evolution. With this end in view we proceed to explore the three well-known types of configurations, the Maclaurin spheroid, the Jacobian ellipsoid, and the binary configuration.

#### 2 (a) *The Maclaurin Spheroid*

Consider a typical Maclaurin spheroid with semi-axes  $a, a, c$ , density  $\rho$  and angular velocity  $\omega$ . Let

$$c = a \cos \gamma, \quad \frac{\omega^2}{2\pi\rho} = \chi, \quad e = \sin \gamma \quad (1)$$

Here  $e$  is the eccentricity of the spheroid,  $\chi$  and  $\gamma$  are parameters connected by the relation\*

$$\chi = \cot \gamma [\gamma (3 + \cot^2 \gamma) - 3 \cot \gamma]$$

For the mass and the angular momentum we have

$$M = \frac{4}{3} \pi a^2 \rho \cos \gamma, \quad \mu = \frac{2}{5} M a^2 \omega \quad (2)$$

Solving (1) and (2) for  $a$ ,  $\omega$ , and  $\rho$  we obtain

$$a = \frac{25}{6} \frac{\mu^2 \cos \gamma}{M^2 \chi}, \quad \omega = \frac{18}{125} \frac{M^{\frac{1}{2}}}{\mu^{\frac{1}{2}}} \left( \frac{\chi}{\cos \gamma} \right)^{\frac{1}{2}}, \quad \rho = \frac{2}{5^{\frac{1}{2}}} \frac{3^{\frac{1}{2}}}{\pi} \frac{M^{10}}{\mu^8 \cos^6 \gamma}$$

\* Thomson and Tait, vol. 2, p. 771 (1883)

The kinetic energy is

$$T = \frac{1}{2} \mu \omega = \frac{9}{125} \frac{M^2}{\mu^2} \left( \frac{\chi}{\cos \gamma} \right)^2$$

The potential energy is

$$W = -\frac{4}{5} \pi \epsilon^2 \rho M = -\frac{18}{125} \frac{M^2}{\mu^2} \frac{\gamma \chi}{\sin \gamma \cos \gamma}$$

We will denote the expressions

$$\left( \frac{\chi}{\cos \gamma} \right)^2, \quad \frac{\gamma \chi}{\sin \gamma \cos \gamma}, \quad \frac{9}{125} \frac{M^2}{\mu^2},$$

by the symbols  $\eta$ ,  $\epsilon$ , and  $A$  respectively. The total energy  $E$  is then  $A(\eta - 2\epsilon)$ . There is one and only one Maclaurin spheroid corresponding to each value of  $\gamma$ , subject to the inequalities  $0 < \gamma < \frac{1}{2}\pi$ . Taking typical values of  $\gamma$  in this region the quantities  $a$ ,  $\epsilon$ ,  $\omega$ ,  $\rho$ ,  $T$ ,  $W$ , and  $E$  are now calculated and the results are set out in Tables I (a), I (b), and IV (a).

It does not appear that the total energy  $E$  has been tabulated up till now, and it is an affair of some trouble, a column for momentum can readily be added to the tables, as it is equal to  $I\omega$ , while  $T$  is  $\frac{1}{2}I\omega^2$ , thus  $E$  can be graphed against density  $\rho$  subject to  $\mu$  constant, and the evolution is towards diminishing  $E$  on this graph.

## 2 (b) The Jacobian Ellipsoid \*

The semi-axes of a Jacobian ellipsoid may be expressed as  $a$ ,  $b = a \cos \beta$ ,  $c = a \cos \gamma$ ,  $\beta$  and  $\gamma$  being subject to  $0 < \beta < \frac{1}{2}\pi$ , also  $54^\circ 21' 27'' < \gamma < \frac{1}{2}\pi$ . A Jacobian ellipsoid does not exist for  $\gamma < 54^\circ 21' 27''$ .  $\beta$  and  $\gamma$  are connected by the relation

$$\sin^2 \gamma \cos^3 \gamma \int_0^\gamma \frac{\tan^2 \phi}{(1 - \kappa^2 \sin^2 \phi)^{3/2}} d\phi = \cos^3 \beta \int_0^\gamma \frac{\sin^2 \phi}{(1 - \kappa^2 \sin^2 \phi)^{3/2}} d\phi,$$

where  $\kappa$  is defined by  $\sin \beta = \kappa \sin \gamma$ .

For the Jacobian ellipsoid to be determinate there must be a relation between  $\chi$ ,  $\beta$ , and  $\gamma$ . It is

$$\frac{\omega^2}{4\pi\rho} = \frac{1}{2}\chi = \cot \beta \cot \gamma \operatorname{cosec}^3 \beta (1 + \cot^2 \beta) (E - E_0) - \frac{\kappa^2}{\kappa^2} E \cot^2 \beta \cot \gamma \operatorname{cosec} \beta + \frac{\cot^3 \beta \cot^2 \gamma}{\kappa^2},$$

where  $\kappa^2 = 1 - \kappa^2$  while

$$E = \int_0^\gamma (1 - \kappa^2 \sin^2 \phi)^{1/2} d\phi \quad \text{and} \quad E_0 = \int_0^\gamma (1 - \kappa^2 \sin^2 \phi)^{-1/2} d\phi.$$

\* Sir G. H. Darwin 'Collected Papers,' vol. 3, p. 123

Since the mass  $M$  and the angular momentum  $\mu$  are assigned we may write

$$M = \frac{4}{3} \pi a^3 \rho \cos \beta \cos \gamma, \quad \mu = \frac{M}{5} a^2 (1 + \cos^2 \beta) \omega,$$

and solving these for  $a$ ,  $\omega$ ,  $\rho$ ,

$$\begin{aligned} a &= \left\{ \frac{5\mu}{M(1 + \cos^2 \beta)} \right\}^{\frac{1}{2}} \frac{2 \cos \beta \cos \gamma}{3M}, \quad \omega = \left\{ \frac{3M\chi}{2 \cos \beta \cos \gamma} \right\}^{\frac{1}{2}} \left[ \frac{H(1 + \cos^2 \beta)}{5\mu} \right]^{\frac{1}{2}}, \\ \rho &= \frac{2}{5\pi} \frac{M^{10}}{\mu^6} \left[ \left( \frac{\chi}{\cos \beta \cos \gamma} \right)^4 \frac{1}{\chi} \left( \frac{1 + \cos^2 \beta}{2} \right)^3 \right], \\ T &= A \left[ \frac{(1 + \cos^2 \beta)^2}{8} \left( \frac{\chi}{\cos \beta \cos \gamma} \right)^3 \right], \\ W &= -A \left[ \frac{(1 + \cos^2 \beta)^2}{4} \frac{2F\chi}{\sin \gamma \cos \gamma \cos \beta} \right] \end{aligned}$$

Defining  $\eta$  and  $\epsilon$  by  $T = A\eta$  and  $W = -2A\epsilon$ , we have  $E = A(\eta - 2\epsilon)$ . Choosing typical values of  $\gamma$ , the quantities  $a$ ,  $\omega$ ,  $\rho$ , etc., are calculated for the corresponding ellipsoids and Tables II and IV (b) are constructed.\*

### 2 (c) The Binary Configuration

The mass  $M$  is supposed to be separated into two detached portions  $M_1$  and  $M_2$  rotating like one connected rigid body. It is convenient to express  $M_1$  and  $M_2$  in the forms  $\frac{M\lambda}{1+\lambda}$  and  $\frac{M}{1+\lambda}$  respectively ( $\lambda \geq 1$ ). Let  $\rho$  be the density, the masses being so far apart that they may be regarded as spherical, with radii  $a_1$  and  $a_2$  and  $r$  the distance between the centres. As the spheres rotate and revolve with the same angular velocity  $\omega$

$$\frac{M}{r^3} = \omega^2 = 2\pi\rho\chi,$$

$$a_1^3 = \frac{3M\lambda}{4\pi\rho(1+\lambda)}, \quad a_2^3 = \frac{3M}{4\pi\rho(1+\lambda)}, \quad \mu = M^{5/3}\omega^{-1/3}\zeta,$$

where

$$\zeta = \frac{2}{5(1+\lambda)^3} \left[ \chi^{2/3} \left( \frac{3}{2} \right)^{2/3} (1 + \lambda^{5/3}) (1 + \lambda)^{1/3} + \frac{5}{2} \lambda \right],$$

$$T = \frac{M^6}{2\mu^3} \zeta^2,$$

$$\begin{aligned} W &= -\frac{4}{5} \pi a_1^2 \rho M_1 - \frac{4}{5} \pi a_2^2 \rho M_2 - \frac{M_1 M_2}{r} \\ &= -\frac{M^6 \zeta^2}{\mu^3} [f_1(\lambda) + \chi^{1/3} f_2(\lambda)], \end{aligned}$$

\* The values of  $F$  and  $E$  needed to complete these computations were taken from Table IX of the second volume of Legendre's "Traité des Fonctions Elliptiques."

where

$$f_1(\lambda) = \frac{\lambda}{(1+\lambda)^2} \quad \text{and} \quad f_2(\lambda) = \frac{2}{5} \left( \frac{3}{2} \right)^{2/3} \frac{1+\lambda^{5/3}}{(1+\lambda)^{5/2}}$$

It is clear that  $\chi$  must be very small for the masses to be nearly spherical under their mutual attraction. An upper limit on  $\chi$  is obtained, taking into account the liability to tidal disruption and partial secular instability. Let us put  $r = a\psi$  where  $a$  is given by  $\frac{3M}{4\pi\rho} = a^3$ . Since  $\frac{M}{r^3} = \frac{M}{a^3\psi^3} = \omega^2$  we have  $\omega^2\psi^3 = \frac{4}{3}\pi\rho$  or  $\chi = \frac{2}{3\psi^3}$ . For the reasons given by Jeans\*  $\psi > 2.5$  roughly. Thus  $\chi < 10^{-2} \times 4/268$ , say,  $\chi_0$ .

A rotating mass cannot break up into two spheres unless  $\chi$  is less than  $\chi_0$ . According to the notation followed so far we may write

$$T = A\eta \quad \text{and} \quad W = -2Az,$$

where

$$\eta = \frac{125}{18} \zeta^3 \quad \text{and} \quad z = \frac{125}{18} \zeta^2 [f_1(\lambda) + \gamma^{1/3} f_2(\lambda)]$$

Let us put

$$B = \frac{M^{10}}{\mu^4} \frac{2}{5^2\pi}, \quad D = \frac{50}{3} \frac{\mu^2}{M^2}$$

Then the numerical values given in the tables for energy, density, and length are  $A^{-1}$ ,  $B^{-1}$ ,  $D^{-1}$  times their actual values respectively. The binary configurations are characterized by two independent parameters  $\lambda$  and  $\chi$  subject to the conditions  $\lambda \geq 1$ ,  $\chi > \chi_0$ . For  $\gamma = \chi_0$  and some values of  $\lambda \geq 1$ , Table III is constructed.

### 3 The Kelvin-Poincaré Evolution

Fig. 1, in which the total energy with its sign changed,  $+E$ ,  $-E$ , is plotted against  $\chi$ , is the diagram of the gradual succession of configurations as  $\gamma$  increases, but the actual evolution is for constant  $\mu$  as *infra*. The course of evolution starts at O, marches along the thick line OA until it is diverted at A to pursue the other thick line up to E. At this point a cataclysm begins to develop which ultimately comes to an end in the neighbourhood of the binary configuration P that has nearly the same energy for the same  $\mu$  and  $\rho$ . The details are as follows —

The line OABCD represents the Maclaurin spheroids, the thick line OA represents the secularly stable spheroids, AC are secularly stable if the figures

\* "Problems of Cosmogony and Stellar Dynamics," p. 54 (1919)



It is natural to expect that in the initial smooth stages of the cataclysm the pear-shaped figure, although unstable, may develop. The following independent analysis, however, indicates that Jeans' figure, unlike the less-developed Darwin-Poincaré "pear," has no place in the evolution.

Since the liquid is not supposed to have become turbulent in such early stages we may consider the density  $\rho$ , the total energy  $E$ , and the angular momentum  $\mu$  to be the same for the critical ellipsoid and the initial forms of the changing pear. In our notation

$$T + W = E, \quad 2T = \mu\omega, \quad \omega^2 = 2\pi\rho\chi, \quad I\omega = \mu,$$

$I$  being the moment of inertia about the axis of rotation. By logarithmic differentiation we obtain for constant  $\mu$ , if  $\rho$  were unchanged,

$$2 \frac{dI}{I} + \frac{d\chi}{\chi} = 0$$

But this is not satisfied by the Jeans' pear (*loc. cit.*, p. 101) for which

$$\frac{d\chi}{\chi} = 0.05227 \epsilon^2, \quad \frac{dI}{I} = -0.09378 \epsilon^2,$$

where  $\epsilon$  is a small parameter, so that it is not in the evolutionary succession.

This shows that the pear-shaped forms cannot arise in the early smooth stages of the cataclysmic transitions unless perhaps some of the implied restrictions relating to constancy of  $\mu$ ,  $\rho$ , and  $E$  are removed. We may examine whether the density slightly increases. Let then the density of the "pear" be  $\rho + d\rho$ ,  $\rho$  being that of the critical ellipsoid. The total mass and the form of the figure remaining the same, the moment of inertia  $I + dI$  is replaced by  $(I + dI)(1 - \frac{2}{3} \frac{d\rho}{\rho})$  which is  $I + dI - \frac{2}{3} \frac{I}{\rho} d\rho$ , to the first order. We have from the conservation of angular momentum

$$I\omega = (I + dI - \frac{2}{3} \frac{I}{\rho} d\rho)(\omega + d\omega),$$

giving

$$\frac{dI}{I} + \frac{d\omega}{\omega} - \frac{2}{3} \frac{d\rho}{\rho}$$

Also as

$$\omega^2 = 2\pi\rho\chi, \quad \frac{d\rho}{\rho} = 2 \frac{d\omega}{\omega} - \frac{d\chi}{\chi}.$$

Thus

$$\frac{1}{3} \frac{d\rho}{\rho} = 2 \frac{dI}{I} + \frac{d\chi}{\chi},$$



so that for Jeans' pear  $d\rho/\rho = -0.40587e^2$ , which means expansion. On the other hand, the formation of the Darwin-Poincaré pear\* demands contraction, which is more natural, since for the latter

$$\frac{d\epsilon}{\gamma} = -0.144306e^2, \quad \frac{dI}{I} = 0.157786e^2,$$

and hence

$$\frac{d\rho}{\rho} = 0.513798e^2$$

If the cataclysm carries over the quantities  $E$ ,  $\rho$ , and  $\mu$  all practically unchanged from the critical ellipsoid, the binary configuration P is reached. The ratio of masses of the pair of bodies corresponding to P is 3.37. The dotted curves in fig. 1 are the equi-density curves, the one passing through P corresponding to the density of the critical ellipsoid. In view of the principle of the degradation of energy, the conclusion seems inevitable that the ratio of masses ( $\lambda$ ) after fission in this model of evolution cannot exceed 3.4. Thus, by the way, is remarkably in agreement with observation for double stars.

Lastly, consider the process of contraction geometrically. If the contraction is due to cooling of the fluid it might perhaps be expected to proceed uniformly in all directions, then every figure should be wholly within the space occupied by the preceding, instead of bulging in parts. Consider first the spheroids.

Let  $r$  be the radius vector to the surface of a Maclaurin spheroid, drawn from the centre, making an angle  $\theta$  with the plane of the equator. Then we have

$$\frac{1}{r^2} = \frac{\cos^2 \theta}{a^2} + \frac{\sin^2 \theta}{c^2}.$$

The contraction  $\delta r$  in the direction ( $\theta$ ) as the matter passes to the consecutive stage is given in terms of the principal contractions  $\delta a$ ,  $\delta c$  by

$$\frac{\delta r}{r^3} = \frac{\delta a \cos^3 \theta}{a^3} + \frac{\delta c \sin^3 \theta}{c^3}.$$

Since

$$a = p \frac{\cos \gamma}{\gamma}, \quad c = p \frac{\cos^3 \gamma}{\gamma},$$

where  $p$  is a constant,

$$-\frac{\delta a}{a^3} = \frac{1}{p^3} \frac{\sin \gamma \delta \gamma + \cos \gamma \delta \chi}{\cos^3 \gamma} \chi, \quad \frac{\delta c}{c^3} = \frac{1}{p^3} \frac{2\chi \sin \gamma \delta \gamma + \cos \gamma \delta \chi}{\cos^3 \gamma} \chi.$$

It can be shown from Tables I (a) and I (b) that  $d\chi/d\gamma > 0$  from  $e = 0$  to

\* Darwin, *loc. cit.*, p. 380

$e = 0.93$  Hence up to this point  $-\frac{\delta a}{a^3} < \frac{\delta c}{c^3}$  Also  $\frac{d}{d\theta}(\delta r) = 0$  when  $\theta = 0$

and  $\frac{1}{2}\pi$  This shows that the minimum contraction is in the equatorial plane and that its value is  $\delta a/a^3$  The maximum expansion is along the axis of rotation and its value is  $\delta c/c^3$   $\delta a$  and  $\delta c$  have been shown to be positive later on All this is quite in keeping with the cooling hypothesis, for the heat generated inside the surface will be conducted more rapidly in the direction of the shortest radius vector, and the subsequent radiation and cooling would produce the maximum contraction in this direction

Table I (a) —Maclaurin Spheroids

$e = \sin \gamma$	$x = \frac{\omega^2}{2\pi\rho}$	$\rho$	$T = \eta$	$e = \frac{1}{2}W$	$2e - \eta = -E$
0.1	$10^{-3} < 2.361$	$10^{-3} < 1.34$	$10^{-3} < 5.63$	$10^{-3} < 2.37$	$10^{-3} < 4.7$
0.15	$10^{-3} < 6.253$	$10^{-3} < 2.56$	$10^{-3} < 4.00$	$10^{-3} < 6.34$	$10^{-3} < 1.26$
0.2	$10^{-3} < 1.159$	$10^{-3} < 1.69$	$10^{-3} < 1.40$	$10^{-3} < 1.19$	$10^{-3} < 2.37$
0.25	$10^{-3} < 1.589$	$10^{-3} < 5.75$	$10^{-3} < 2.69$	$10^{-3} < 1.65$	$10^{-3} < 3.29$
0.3	$10^{-3} < 2.49$	$10^{-3} < 1.73$	$10^{-3} < 6.47$	$10^{-3} < 2.588$	$10^{-3} < 5.111$
0.35	$10^{-3} < 3.315$	$10^{-3} < 4.73$	$10^{-3} < 1.25$	$10^{-3} < 3.61$	$10^{-3} < 7.11$
0.4	$10^{-3} < 4.356$	$10^{-3} < 1.17$	$10^{-3} < 2.25$	$10^{-3} < 4.89$	$10^{-3} < 9.66$
0.5	$10^{-3} < 6.90$	$10^{-3} < 5.84$	$10^{-3} < 6.34$	$10^{-3} < 8.37$	$10^{-3} < 1.61$
0.6	$10^{-3} < 1.007$	$10^{-3} < 2.49$	$10^{-3} < 1.58$	$10^{-3} < 1.35$	$10^{-3} < 2.54$
0.7	$10^{-3} < 1.387$	$10^{-3} < 1.027$	$10^{-3} < 3.77$	$10^{-3} < 2.15$	$10^{-3} < 3.93$
0.75	$10^{-3} < 1.597$	$10^{-3} < 2.13$	$10^{-3} < 5.83$	$10^{-3} < 2.73$	$10^{-3} < 4.88$
0.8	$10^{-3} < 1.812$	$10^{-3} < 4.58$	$10^{-3} < 9.117$	$10^{-3} < 3.499$	$10^{-3} < 6.088$

Table I (b) —Maclaurin Spheroids

$e = \sin \gamma$	$x = \frac{\omega^2}{2\pi\rho}$	$\rho$	$T = \eta$	$e = \frac{1}{2}W$	$2e - \eta = -E$
0.81	0.18597	$10^{-3} < 5.43$	0.1005	0.3090	0.6387
0.81267*	0.18712	$10^{-3} < 5.678$	0.1031	0.3748	0.6465
0.82	0.19039	$10^{-3} < 6.42$	0.1103	0.3898	0.6990
0.85	0.2029	0.1085	0.1484	0.4602	0.7721
0.9	0.2203	0.296	0.2556	0.6290	1.0024
0.91	0.2225	0.372	0.2890	0.6740	1.0800
0.92	0.2241	0.477	0.3275	0.7225	1.1175
0.93	0.2247	0.621	0.3736	0.7857	1.1974
0.94	0.2259	0.827	0.4309	0.8530	1.2737
0.95	0.2219	1.139	0.5021	0.9340	1.3659
0.96	0.2160	1.636	0.5945	1.032	1.4695
0.97	0.2068	2.518	0.7209	1.158	1.5961
0.98	0.1890	4.295	0.9013	1.323	1.7447
0.99	0.1551	9.44	1.21	1.587	1.9650
1	0	$\infty$	$\left(\frac{\pi^2}{4} - \right) 2.435$	$\left(\frac{\pi^2}{4} - \right) 2.435$	$\left(\frac{\pi^2}{4} - \right) 2.435$

\* Critical spheroid

Table II — Jacobian Ellipsoids

$\epsilon = \sin \gamma$	$\gamma$	$\beta$	$\chi = \frac{a^2}{2\pi p}$	$\rho$	$T = \eta$	$\epsilon = \frac{1}{2}W$	$2\epsilon - \eta = E$
	$^{\circ}$ $'$ $''$	$^{\circ}$ $'$ $''$					
0 81267	54 21 27	0 0 0	0 18712	$10^{-2} \times 5.678$	0 1031	0 3748	0 6465
0 81915	55 0 0	14 15 0	0 187	$10^{-2} \times 5.6908$	0 10315	0 3751	0 6471
0 8358	57 0 0	28 30 0	0 186	$10^{-2} \times 5.935$	0 1052	0 3798	0 6545
0 866	60 0 0	40 54 0	0 18120	$10^{-2} \times 6.858$	0 1115	0 3964	0 6814
0 9064	65 0 0	54 46 0	0 16590	$10^{-2} \times 1.133$	0 1371	0 4599	0 7828
0 93859*	69 49 0	64 23 40	0 14200	0 253	0 18948	0 5817	0 9739
0 9397	70 0 0	64 44 0	0 14094	0 2646	0 1932	0 5900	0 9868
0 9659	75 0 0	72 36 0	0 1072	0 9	0 3105	0 835	1 3605
0 984	80 0 0	79 10 0	0 06614	4 899	0 5691	1 664	2 7589
0 9962	85 0 0	84 52 0	0 02580	76 45	1 404	2 033	3 862
1	90 0 0	90 0 0	0	$\infty$	$\infty$	$\infty$	$\infty$

\* Critical ellipsoid

Table III — Double Figure Configurations

$\chi = \chi_0$				
$\chi$	$\rho$	$T = \eta$	$\epsilon$	$2\epsilon - \eta = E$
1	0 699	0 1698	0 6980	1 228
2	0 3913	0 1292	0 6046	1 08
3	0 183	$10^{-2} \times 8.834$	0 4780	0 8688
4	0 093	$10^{-2} \times 6.297$	0 3915	0 7200
5	0 05108	$10^{-2} \times 4.709$	0 3287	0 6103
6	0 03188	$10^{-2} \times 3.861$	0 2841	0 5297
7	0 02088	$10^{-2} \times 2.985$	0 2490	0 4681
8	0 0145	$10^{-2} \times 2.487$	0 2226	0 4204
9	0 01055	$10^{-2} \times 2.122$	0 2018	0 3824
10	0 007951	$10^{-2} \times 1.842$	0 1848	0 3512
100	$10^{-2} \times 1.678$	$10^{-2} \times 2.075$	$10^{-2} \times 5.458$	0 1065
1000	$10^{-2} \times 8.374$	$10^{-2} \times 1.889$	$10^{-2} \times 4.367$	0 08545
$\infty$	$10^{-2} \times 7.771$	$10^{-2} \times 1.82$	$10^{-2} \times 4.264$	0 08351

In the preceding we took for granted that  $\delta a$ ,  $\delta c$ , the contractions in  $a$ ,  $c$  respectively, are positive. Table IV (a) shows that this is actually so. Table IV (b) for the axes of ellipsoids shows that  $b$  and  $c$  decrease so that  $b/c$  tends to unity as the material contracts. On the other hand,  $a$  gradually increases from 0.7785 to 0.845 and then decreases to nothing. Such a gradual increase and decrease might tend to produce heterogeneity, but we suppose that there is time for the densities to be equalized.

Lastly, I should like to express my thanks to Sir J. Larmor, who pointed out the historical interest of the problem and made many valuable suggestions and criticisms while this work was being carried out.

Table IV (a) —Maclaurin Spheroids

$e$	$a = b$	$c$	$e$	$a = b$	$c$
0 1	105 3	105	0 81267	0 7785	0 4536
0 15	39 54	39 05	0 85	0 649	0 3419
0 2	21 11	20 65	0 9	0 4046	0 2360
0 25	15 23	14 71	0 91	0 4657	0 1931
0 3	9 81	9 37	0 92	0 4368	0 1713
0 35	7 03	6 61	0 93	0 409	0 150
0 4	5 26	4 82	0 14	0 3811	0 130
0 5	3 14	2 72	0 95	0 3528	0 1077
0 6	1 99	1 59	0 96	0 3243	0 09084
0 7	1 36	0 92	0 97	0 2945	0 07155
0 75	1 04	0 684	0 98	0 2633	0 05244
0 8	0 8280	0 4968	0 99	0 2372	0 03305
0 81	0 7887	0 4623	1	0 1462	0

Table IV (b) —Jacobiian Ellipsoids

$e = \sin \gamma$	$e' = \sin \beta$	$e'' = \frac{\sqrt{b^2 - c^2}}{b}$	$a$	$b$	$c$
0 81267*	0	0 81267	0 7785	0 7785	0 4536
0 819	0 246	0 806	0 7905	0 766	0 4534
0 839	0 478	0 784	0 819	0 720	0 446
0 8600	0 6547	0 7500	0 845	0 638	0 422
0 9063	0 8168	0 6807	0 827	0 477	0 349
0 93859†	0 90183	0 6024	0 7454	0 3221	0 2572
0 9397	0 9042	0 5991	0 7369	0 3150	0 253
0 9659	0 9542	0 5014	0 6077	0 1817	0 1573
0 9648	0 9822	0 3822	0 4806	0 0886	0 080
0 9962	0 9960	0 2313	0 2970	0 02658	0 02589
1	1	0	0	0	0

\* Critical spheroid

† Critical ellipsoid

Addison December 18, 1933

*"On Astronomical Evolution by Dissipation of Dynamical Energy"*

By SIR JOSEPH LARMOR

The object of the analysis in this paper, as stated in the beginning, is to set out in more systematically developed view the Kelvin succession of events in his historical problem, and especially to illustrate the general theory of dynamical evolution. For the dynamics of the planetary system, now so accurately developed, is concerned with the residual cyclic motions, which persist after non-cyclic features have been smoothed away in course of ages

by frictional forces, tidal, and other influences. In the present illustration it is the trend of an energy-diagram that determines the course of the evolution.

In a material system in pure dynamics the variables are a set of co-ordinates typified by  $q$  with their component velocities  $\dot{q}$  and momenta  $p$  which are  $\partial T/\partial \dot{q}$ . Here  $T$  is the kinetic energy which is a function of the variables  $\dot{q}$  with coefficients functions of  $q$ ; there is a potential energy  $W$  so that  $T + W$  is total energy which as was known in early days, is conserved in an isolated frictionless system. But of these variables there may be a set of co-ordinates of type  $\psi$  which do not appear in the expressions for  $T$  and  $W$  with velocities  $\dot{\psi}$  and momenta  $\partial T/\partial \dot{\psi}$  or  $\Psi$ . These momenta are conserved being cyclic in type, and the velocities related to them can be eliminated from  $T$  by means of the set of equations  $\partial T/\partial \dot{\psi} = \Psi$ . The Lagrangian generalized dynamics, as developed into cyclic co-ordinates by Routh, Kelvin, and Helmholtz replaces the Lagrangian function  $L = T - W$  by a modified function  $L'$  which is  $\tau_2 + \tau_1 + \tau_0$  where the terms are of the second degree, first degree, and degree null, in the remaining velocities of type  $\dot{q}$ , with coefficients involving the  $q$  and also the  $\Psi$  which are now constants of the motion. The equations of motion are then of the form

$$\frac{d}{dt} \left( \frac{\partial L'}{\partial \dot{q}} \right) - \frac{\partial L'}{\partial q} = F_q,$$

where  $F_q$  is the extraneous force, in addition to the force involved in  $W$ , acting on the co-ordinate  $q$ . The time-rate of gain of energy of the system is  $\Sigma F_q \dot{q}$  which works out into the complete differential expression  $dE/dt$  where  $E = \tau_2 + \tau_0$ , in which the first order terms  $\tau_1$  which alone would change sign on reversal of the action even though the cyclic momenta have not been reversed have disappeared. The expression  $L'$  is equal to  $T - W - \Sigma \dot{\psi} \partial T/\partial \dot{\psi}$  in which the variables  $\dot{\psi}$  have been eliminated by introducing the constants  $\Psi$  instead of them. All the equations of motion are expressed from  $L'$  alone and it might be hastily presumed that the course of evolution by dissipation would depend on the change of the time-gradient of the effective Action, which is  $L'$ . But that is not so, although  $L'$  by its  $n$  equations determines the course of motion completely. The energy that comes into the system is the change in  $E$ , and it is  $E$  that is subject to dissipation within the system by frictional agencies not involved in  $W$ . Then the course of a steady evolution, as in the discussion above is to be presented as one of continual fall of  $E$  subject to conserved cyclic momenta. If the co-ordinates are cyclic so that  $\dot{q}$  are not involved in the energy the form of  $E$  is static varying with the co-ordinates  $q$  alone, and the steady so-called secular states of the system are those for which

$E$  is minimal. We can represent the course of this evolution graphically. Let the variables  $q$  represent Cartesian co-ordinates in a hyperplane of the same number of dimensions, and imagine an ordinate  $E$  drawn at each point of it. A set of hypersurfaces each of constant energy  $E$  is obtained, and the bottom points represent the steady states of the system. The hypersurfaces of constant  $E$  will now take successive forms owing to change by internal dissipation, in simple cases changing an internal parameter. On this succession of hypersurfaces there will be continuous sets of lowest points which trace out curves of smooth evolution. The highest points and cols of saddle shape will represent unstable positions, impermanent but of change initially slow. It is very unlikely that two of these curves of evolution will meet and cross each other, unless there is one co-ordinate  $q$ , when the space is a plane with co-ordinates  $E, q$ . A general principle was advanced by Poincaré on graphical grounds that when a curve of stability meets and crosses one of the other set of curves, those of slowly changing unstable forms, their condition as to stability interchanges, this is the normal course of crossing, for two curves of stability cannot come together unless by way of a curve of summits of the other kind which lies between them. The principle predicted that the pyriform figure is stable but the result now seems to be accepted, after Jeans, that the pyriform states are initially unstable, in opposition to the Poincaré\* criterion. This may be because a real crossing point is very seldom reached, as the number of independent variables for a fluid mass is not finite, for Poincaré himself appears to arrive at an infinite series of transitions (cf Lamb's account) by a Lamé analysis of the normal states of vibration of the fluid mass. Thus the mode of lapse into static states is determined by a set of equations equal in number to the effective co-ordinates, of which the equation of energy is only one, yet one which is involved along with all the others in the Action Principle and its characteristic function  $L'$ . Our evolutionary criterion is far less complete, is merely one for the gradual rough trend of steady evolutionary states, without detail as to the transition which remains unexplored. This general criterion of trend, the only one there is, prescribes total energy falling away for conserved cyclic momenta.

\* Correction of Poincaré's original investigation (1885) was soon made by Schwarzschild in a discussion which is difficult of access. A very complete historical account was given at an early stage by H. Lamb, 'Hydrodynamics,' ed 1895 and ed 1932, where he reproduces some numerical tables from Kelvin and Darwin which ought to agree with those worked out and applied above. There is an account by von Ziepel in the Poincaré Memorial Volume of "Acta Mathematica." A formal treatise by L. Lichtenstein is announced which may provide closer detail.

But the main point is to assert the essential limitation of most of these evolutionary discussions. The subject originated with W. Thomson's formulation of the Dissipation of Energy on the basis of the formal Carnot doctrine, at first without the illustrations from statistical atomic theory which made it so natural. He was soon deflected from the development of his thermodynamic doctrine of Available Energy by the obvious problems of dissipation presented on a vast scale by astronomical systems. If with Kelvin, and Helmholtz, and Waterston even earlier, the availability that is inevitably dissipated is taken to be in part the energy of gravitational collocation of the bodies concerned, a problem is presented of which a very striking analytical exemplification has been prepared gradually by the elegant mathematical work of Jacobi and Kelvin and Poincaré. But this criterion of dissipation confined to energy purely mechanical is only valid, on the Carnot-Kelvin doctrine, under limitation of the system to uniformity and constancy of temperature and of state during its evolution. If the state is liable to change but restricted to constant temperature, it is a different entity, Thomson's Available Energy as distinct from total energy, that is the subject of dissipation. If the temperature can change also, the formulation of a criterion is more complex. It is the special merit of the Clausius procedure by introduction of an Entropy inherent in the system—an offshoot from the Carnot-Kelvin temperature theory, which was later put into illustrative relation with the doctrine of probabilities of molecular states, and has now become a foundation postulate in its own right in these quantal days, specified as an assumed definite function of state and configuration and temperature for every system—that it has covered the ground in a more direct manner, because Entropy trends upward without limitation except such as arises from abstraction of the rather indefinite entity experienced as heat. Entropy may be transferred, but cannot diminish *as a whole*. Any precise definition of heat must indeed be by way of Entropy or of Available Energy. Thus for a gaseous star subject to its immense changes of temperature on shrinkage, the gravitational energy by itself has little to do with the secular course of evolution, except as the merest illustration. In the abstract hydrodynamical problem that fascinated Jacobi, Helmholtz, Poincaré, mainly on Kelvin's initiative, it is, however, reduced to an affair of structureless fluid, of unchanging density and therefore also unchanging temperature, so that as it happens the criterion of falling gravitational energy does apply, as indeed under such limitations it could not help doing. If these conditions were not satisfied, though abundance of types of nearly steady state might be worked out, and their ranges of dynamical stability marked,

there would be no practicable general criterion, so long as discussion by Entropy or Available Energy is unmanageable, of trend of change as in fact governed by dissipation of availability, to decide how far the crowd of possible solutions even belonged to the same linear evolutionary succession, except by presumptions derived from comparisons regarding the ordinary dynamical stabilities of the separate states that have been usually employed. Following Poincaré it was commonly held that these questions, which belong really to a pure *topology* of the various linear series of steady secular states and the transitions where they meet, could be readily settled by a principle of exchange of stabilities, though it appears somewhat mysterious that a consideration of linear topological grouping alone should settle a definite dynamical stability where there may be many degrees of freedom involved, obvious or latent, indeed, for the special example now on hand, in the transition from the Jacobian to the pyriform secular figures, actual discrepancy appears to have been encountered. A final remark may not be superfluous, that this doctrine of dissipation, which is wider than any atomic scheme, does not stand in any relation to the principle of Least Action, which belongs to reversible dynamics, though Boltzmann, Clausius, Burbury, and others at one period tried hard to connect it up with the foundations of the principle of Carnot.

The feature that has invited close attention to the special Kelvin problem of evolution is its asymmetric character. Starting with the Maclaurin oblate spheroids, becoming more oblate as they lose energy ( $E$ ) while momentum is conserved, the series continues as Jacobian ellipsoids with loss of the axial symmetry, and then, as is suggested, it passes on into lopsided forms, the pyriform bodies of Poincaré and Darwin, which may gradually develop a constriction and finally separate into two masses of the type of the actual double stars. Without that hint from the heavens such irregular forms might hardly have been thought of as arising in the course of evolution. One notes that the conserved momentum  $G$  being  $I\omega$ , can readily be inserted in the tables. The total energy  $E$  does not appear to have been previously examined at all. The Maclaurin stability is found to persist, but only for axial disturbance, beyond the point of bifurcation into Jacobian forms, up to the limit determined by G. H. Bryan. Possible initiation towards a nebular ring-structure is perhaps also not quite improbable, of the astronomical phenomena of *Novæ*, the mathematical theory was explored long ago (cf. Lamb, § 376) showing as is natural that for a ring of fluid type the period of rotation would be about the same as for meteoric bodies such as are required for stability of



the Saturnian rings. The tendency to such incipient ring-structure of compressible non-uniform types may well be held to have some persistence before breaking up into parts flying off tangentially, in arrangement more or less symmetrical with regard to the axis, after the manner of the wings of the spiral nebulae. The observed rotational velocities  $v$ , of the Andromeda nebula, are not inconsistent with such an origin. The fundamental contrast with the very peculiar but dynamically well-founded lopsided type of evolution towards a double star is connected by Jeans ("Cosmogony") with this expansile tendency of the outer parts.

The subject matter of this discussion developed itself some years ago for class lectures at Cambridge on the general theory of Energy. The discrepancy of stabilities, as independently determined, with the Poincaré criteria has persisted. The Poincaré series of configuration (cf. Lamb's discussion, § 377) are series with changing  $G$ , the series of an actual evolution are of  $G$  necessarily constant but changing  $E$ . The argument has regard on one side to unrestricted continuous change expressible in terms of differentials of first and second orders, so that stabilities as regards the various co-ordinates can be discussed for each energy state independently of one another, in contrast to the turbulent transitions, necessarily largely undetermined and mixed up, in the path of secular evolution. The suspicion here arising would in effect be whether the use of the same differential symbol in discussing ordinary dynamical stability on a given energy-level, and in representing the undetermined frictional transition to the other energy surfaces, is not analogous to the ancient ambiguity between differentials and variations in the variational calculus, which was brought out and surmounted by the notation and procedure of Lagrange.

A very remarkable actual fact may here be relevant. An arresting feature of the astronomical landscape is the presence of open groups of stars, all of them, though far apart, with the same velocity towards the same direction, in the case of the *Ursa Major* cluster the Sun happens to lie within the group, so that its stars appear all over the sky. These groups cannot have been for long time within our stellar system, else their equality of velocities would have been deranged. They must have come in from the outside more empty space, and are perhaps merely passing across—though our own system is itself presumably scattering. The stars of each group travel together like a solid body, but as one which has acquired no rotation comparable with its translatory motion. The problem of the origin of the group thus arises. A solution that suggests itself is that the group has broken off from a much larger

cosmic system, for which rotational motion as a whole would be translatory motion for a local group in it. For example, it may have been shed off from an arm of a spiral nebula, when it was in position on the nebula internal gravitational disturbance was compensated, while after it has broken off, on its journey across space with its initial velocity, it need not have had time to cumulate much disturbance. The group could not have come from a very distant nebula for yet another reason: its velocity would be excessive and in the wrong direction. If this be tenable, the velocities being of suitable order, the empty cosmic spaces would be inhabited by such relatively small isolated stellar groups as well as by isolated single stars.

A general key is indicated above in the sharp distinction between differentials and variations, introduced into analysis by Lagrange, which made general dynamics a possibility in connexion with his Calculus of Variations. For a given form of the energy function  $E$  the stabilities of the system may be explored by the *differential* dynamical theory, adjacent forms of  $E$  differ by the coefficients in its analytic form having received *variational* change. In general there would be as many types of variation as there are co-ordinate variables. But if we assert limitation of the modes of variation to one kind, expressed by one parameter  $\lambda$  (cf. Lamb, § 377) we can draw a series of graphs of  $E$  as a function of  $\lambda$  alone, thus of two-dimensional character, and the Poincaré theory of intersections with interchange of stabilities acquires a footing. But this limited variation is hard to conceive in general and makes a large call on previous special knowledge of the system. Such knowledge is available in the very special problem of rotating perfect fluid. It recognizes the actual existence of a plurality of states, while it has to *postulate* that they are not a mere unrelated assemblage of self-subsisting forms, but express an actual mode of progress in the evolution of the system as a linear series. This means that it will not branch off into other modes except where this one becomes unstable, and such ordinary instability is sufficiently explored by neglecting slow dissipation, in the hydrodynamic problem by neglecting viscous terms in the equations. Here the Jacobi series is an example: they are stable from their beginning as a branch from the Maclaurin series until a place where a pyriform series branches off, it was found by Poincaré (Lamb, § 180) that afterwards complete change of character arises, that there are an infinite number of bifurcations each originating a branch series, a remarkable transition which perhaps derives from the abstract simplicity of ideal fluid systems.

The general theory gets a foothold only when we have an assurance in advance that there is a definite linear order of succession of states, that the evolutionary variation of  $E$  depends (somehow) on only one parameter  $\lambda$ . In the Kelvin hydrodynamic problem the restriction that is tacitly imposed, involving this result, is that in each stage the system is to be moving like a solid body, implying an evolution of extreme slowness, the general possibilities of freedom would be far wider than this

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*The Emission of Electrons under the Influence of Chemical Action*  
*Part II —Some General Conclusions and a further Study of the*  
*Case of Carbonyl Chloride*

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§1 *Some General Conclusions*

We have now investigated this effect on the sodium potassium alloy  $K_2Na$  for 22 different gases. With 15 of these the emission has been found to be sufficiently large for the distribution of kinetic energy among the emitted electrons to be determined, using the reacting gases at pressures of the order of  $10^{-5}$  mm of mercury.

As a result of these experiments we have been led to some general conclusions, the most important of which we propose to state, very briefly, at the outset. They are —

(1) The energy distribution is not in general of the Maxwellian type as the earlier evidence had tended to indicate. We expressed a suspicion of the correctness of this view on p 49 of our previous paper\*. The later experiments have confirmed that suspicion.

(2) The distribution curves all rise from a small value at zero energy ( $V = 0$ ) to a maximum at a certain energy ( $V = V_m$ ) and fall away to zero as the energy is increased further to  $V = E_a$ . In this statement zero is to be understood to mean zero within the limits of our experimental error.

\* 'Proc Roy Soc,' A, vol 132, p 22 (1931). This paper contains references to earlier publications.

(3) The *actual* maximum energy  $E_a$  of the emitted electrons is ultimately reached rather gradually and is difficult to determine precisely. The experimental plots, in fact, show a 'tail' similar to that observed in the photoelectric effect. There is a *practical* maximum energy  $E_m$  obtained by disregarding the tail, 99.9% of the emitted electrons have energies less than  $E_m$  in the case of  $\text{COCl}_2$ .

(4) The distribution curves for energies  $> V_m$  and  $< E_m$  can all be represented quite closely by the equation

$$N(V) dV = Ae^{-k(V - V_m)^c} dV, \quad (1)$$

where  $A$ ,  $k$ , and  $c$  are constants. The constant  $c$  has a value very close to 2 for the more energetic and very close to 1 for the less energetic reactions. For the most energetic reactions the equation (1) covers approximately the whole range of emitted electrons from  $V = 0$  to  $V = E_m$ , for these reactions the distribution curves, if we disregard the 'tails,' are almost symmetrical about  $V = V_m$ .

Equation (1) is inconsistent with the existence of a maximum energy  $E_a$  such as is referred to under (2). It should be remembered, however, that equation (1) is not a theoretical equation. It is a convenient empirical way of representing some of the facts.

(5) For chlorine compounds (for other substances we have not sufficient data) there is a simple relation between the maximum energy  $E_m$  of the emitted electrons and the energy of dissociation  $D$  of the chlorine compound referring to the corresponding reaction responsible for  $E_m$ . It is

$$E_m + D = \text{constant} \quad (2)$$

(6) At low pressures the electron emission is proportional to the pressure of the reacting gas. In most gases it rises to a very sharp maximum and then falls, as has been described in Part I for  $\text{COCl}_2$ . However, this is not completely general, in  $\text{COS}$ , for example, the rate of rise of emission with increasing pressure steadily diminishes to zero, so that the emission at higher pressures reaches a constant value independent of the pressure.

(7) If we define the electronic yield  $Y$  of the reaction as the limiting value at  $p = 0$  of the emission divided by the pressure  $p$ , the yield is greatest for the more energetic and least for the less energetic reactions. For  $\text{Cl}_2$ , for example, the maximum number of electrons emitted for each chlorine atom taking part in the reaction is about  $1.5 \times 10^4$  whereas for  $\text{SOCl}_2$  the corresponding

quantity is about  $1.5 \times 10^{-6}$ . The contraction in the yield is very large compared with the contraction in the chemical energy  $E_s$  (see below), as this quantity diminishes. If we plot  $\log Y$  against  $E_s$ ,<sup>1</sup> the points tend to group themselves about a straight line.

(8) A fair account of the facts, at any rate so far as the reactions which involve the formation of alkali-halogen bonds are concerned, can be derived from the following hypothesis: as a result of the collision of a molecule of the chemically active gas with the metal surface a stable polar chemical bond is formed. At the instant of formation the bond consists of a free negative ion, which is one of the atoms of the gas molecule, and one of the positive ions of the metal, which is chemically bound to other atoms of the metal. The kinetics of the heterogeneous change for polyatomic gas molecules may either be that the two active atoms of the gas molecule simultaneously enter into the reaction with the metal, or it may be that only one atom enters into the reaction at the instant of collision with the metal, it depends upon the strength of the bonds between the atoms of the gas molecule and upon the orientation of the molecule at the instant of collision. As we are concerned with the formation not of a diatomic alkali-halogen molecule (or alkali-pseudohalogen molecule) but of only one chemical bond of a large cluster, for example,  $\text{Cl}^- - \text{K}^+ - \text{Rest}$  of the metal, there will be no quantum restrictions for the stabilization of the nascent bond. The stabilization of the bond may occur by the excess energy being dissipated among a number of the adjacent metal atoms by the ordinary mechanism of the heat conductivity of metals. The energy of the reaction will be thus transformed into heat. However, since in the chemical process of formation of a *polar bond* from neutral atoms, or molecules, a *spontaneous re-arrangement of the electronic system* of the nascent bond is involved, the stabilization of the bond may also occur spontaneously under the mechanism of a three-body collision with one of the free metallic electrons. In this case all the energy of the reaction goes off as kinetic energy of the electron which is thus enabled to escape from the metal surface. These escaping electrons constitute the electronic spectrum which is the principal subject of the investigation.

On the basis of this hypothesis we can give an explanation of the empirical energy distribution function given by equation (1). The analysis shows that the observed form of the distribution function can be qualitatively accounted for by the form of the function expressing the probability of the collisions of the second kind for electrons having different velocities (the de-excitation function).

Furthermore, on the basis of this hypothesis, relation (2) receives a very simple interpretation. It is

$$E_m = E_0 - \phi, \quad (3)$$

where  $E_0$  (cf § 6) is the energy of the corresponding elementary chemical reaction responsible for  $E_m$ , and  $\phi$  is the work function of the metal. This is true within the limits of accuracy of our estimation of  $E_0$ . This relation is general, at any rate so far as we have been able to find the data to test it. *The equation (3) is similar to the fundamental law of photoelectricity.*

The evidence which has led us to these conclusions, as well as to some others which will appear in due course, is set forth in this paper and others which will follow it.

Much of this paper will be devoted to an account of further experiments with  $\text{COCl}_2$ . This gas has been chosen for a particularly detailed investigation because it gives an emission of a suitable magnitude, it is convenient to experiment with and we have had more experience in manipulating it than the other gases to be dealt with later.

## § 2 *The High Energy Part of the Electronic Spectrum from Carbonyl Chloride*

As we have stated already, the results of the experiments described in Part I indicated a possible deviation from the Maxwell distribution of energy for the emitted electrons with high velocities. Further consideration led us to attach much more importance to this point, particularly in view of the great deviation from the Maxwell distribution in the region of low velocities which had been observed in all the previous researches. We therefore decided to investigate this question carefully and in detail.

We have to bear in mind that for the determination of the energy distribution among the electrons, the characteristic curves—electron current  $i$ , as a function of the retarding field  $V$ , were employed. For a Maxwell distribution

$$i = i_0 (1 + \sigma V) e^{-\sigma V}, \quad (4)$$

where  $i_0$  is the electron saturation current, and  $\sigma = e/kT$ . This relation held, within the limits of experimental error, in the range of values  $i/i_0 \approx 0.1$  to  $i/i_0 \approx 5 \times 10^{-4}$ . Commencing with  $i/i_0 \approx 0.1$  and going up to  $i/i_0 = 1$ , a great deviation from the above relation was observed. In view of the small values of the electron current at large retarding fields, and consequently

large relative error of measurement, determinations were made only down to  $i/i_0 \approx 5 \times 10^{-4}$

The purpose of the present experiment consisted mainly in the determination of the energy distribution among the electrons in the region of higher velocities. Taking into consideration the experience obtained from the previous work, it was possible to improve upon the accuracy in the measurement of the electron currents, and by such means it was possible to increase the range of measurements and reach a value of  $i/i_0$  so low as  $1 \times 10^{-6}$ . For this, it was essential that the following conditions should be fulfilled —

- (1) Sufficiently high electron saturation current. This limits the range of gas pressure within which the determinations can be made.
- (2) High constancy of gas pressure during the whole period of the experiment. In Part I it was established that the contact potential difference was a function of the gas pressure, and it was the inconstancy of the gas pressure which mainly determined the error of measurements in obtaining the characteristic curves.
- (3) It was necessary to add a suitable system of capacitances, connected in parallel with the electrometer, which would make it possible to vary the sensitivity of the measuring apparatus quickly.

Determinations were made on the same alloy, of the composition approximately  $K_2Na$ , as in Part I and with the same sample of phosgene. The description of the apparatus, and the procedure used in the measurements are the same as there described.

The previous work on the electrons in the range  $i/i_0 = 5 \times 10^{-4}$  to  $i/i_0 = 0.1$  had led to the conclusion that the distribution of energy among the electrons follows Maxwell's law. The average energy corresponding to such a distribution was determined\* by the use of a graph expressing the relation between  $\sigma V$  and  $V_1$ , where  $\sigma V$  is taken from the equation  $i = i_0 (1 + \sigma V) e^{-\sigma V}$ , and  $V_1$  is the experimental scale of potential  $V = V_1 + K$ , where  $K$  is the contact potential difference between the alloy and the outer electrode. For a Maxwell distribution this graph is a straight line, the slope of which is equal to  $\sigma = e/kT$ . On such a graph the deviations from the Maxwell distributions are easily seen as a departure from rectilinearity.

Five characteristic curves  $i/i_0 = f(V_1)$ , obtained under different phosgene pressures from  $1.5 \times 10^{-3}$  mm. to  $3 \times 10^{-4}$  mm., served as data for the

\* O. W. Richardson and M. Brotherton, 'Proc. Roy. Soc.,' A, vol. 115, p. 20 (1927).

present analysis The maximum of electron saturation current  $i_0$  is at about  $5 \times 10^{-5}$  mm, at a pressure of  $1.5 \times 10^{-6}$  mm,  $i_0$  was about 25% and at  $3 \times 10^{-4}$  mm about 50% of the maximum saturation current Thus the observations were made in the most interesting range of pressures, where the important changes in the surface conditions of the alloy occur A further decrease, or increase, in pressure would inevitably have led to a further decrease of saturation current, and in consequence the relative accuracy of the measurements would have been insufficient

The time  $T$  of the drop in all experiments was 25–26 seconds The time of exposure  $dt = 25$  seconds The maximum diameter of the drop was about 4 mm and the drops had a regular spherical shape The maximum saturation current under such conditions was equal to  $(i_0)_{\max} = 1.70 \times 10^{-8}$  amp

Figs 1 and 2 show the above-mentioned graphs  $\sigma V = F(V_1)$

The curves III, V, and II, fig 1, are shifted to the right relative to curve I by 0.3, 0.8, and 1.8 volts respectively The experimental points for curve I are indicated by crosses, the others by circles Table I shows separately the results for one of the curves (curve IV, fig 2) For the purpose of comparison with the results of the previous experiments, a dotted straight line is drawn for each curve This line is so placed that the points of the curve in the range of values of  $\sigma V$  from 3 to 8 would have given the best coincidence with the dotted line The slope of this straight line,  $\sigma$ , has a value of about 5.5, i.e., practically the same value as in the previous determinations Generally, the curves of the previous research, which have been obtained in the range of  $\sigma V$  between 3 and 8, completely coincide with the present curves.

We wish to draw attention to the following features of these curves —

- (1) The great similarity of all the curves among themselves Within the limits of experimental error they coincide with one another
- (2) The slope of the curves continuously increases as  $\sigma V$  increases This can also be seen from Table I In column 3 are given the ratios of each preceding value ( $i'$ ) of the current to the one following ( $i$ ), for every 0.1 volt With an increase in the retarding field ( $V_1$ ) the ratio  $i'/i$  systematically increases The slope, which has the value about 5.5 has a local character and has been obtained because the observations were limited to values of  $\sigma V$  below about 8

From this we arrive at the following important conclusion *The fundamental distribution of energy among the electrons is not according to Maxwell's law*



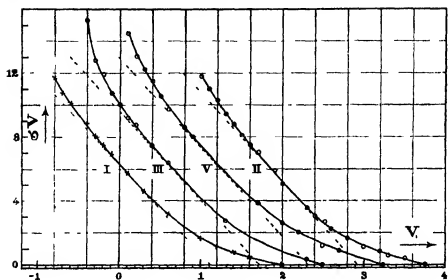


FIG 1

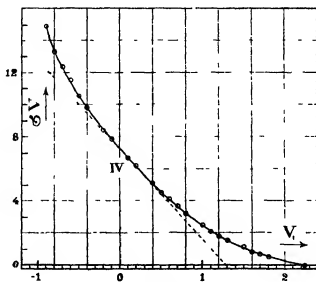


FIG 2

- Curve I .  $p = 2.2 \times 10^{-5}$  mm,  $i_0 = 64 \times 10^{-10}$  amp  
 Curve II  $p = 1.4 \times 10^{-5}$  mm,  $i_0 = 42 \times 10^{-10}$  amp  
 Curve III  $p = 2.0 \times 10^{-5}$  mm,  $i_0 = 102 \times 10^{-10}$  amp.  
 Curve IV  $p = 5.0 \times 10^{-5}$  mm,  $i_0 = 138 \times 10^{-10}$  amp  
 Curve V  $p = 2.7 \times 10^{-4}$  mm,  $i_0 = 36 \times 10^{-10}$  amp

The above evidently solves the question as to the reality of the deviations from the linear relationship in the region of low velocities, which have been observed in all previous researches. We can see now that the results of the previous experiments also indicated deviations from the linear relationship in the region of high velocities. Such deviations can be seen, for example, in fig. 10, p 48 of Part I, and also in the paper by Richardson and Brotherton (*loc cit*, pp 25-26, Table IV and fig 2)

Table I

Volts $V_1$ (CPD = -2.24)	$v/v_0$	(at $V_1' - V_1 = 0.1$ )	$\sigma V$
+2.1	0.954	-	0.338
2.0	0.933	1.02	0.431
1.9	—	—	—
1.8	0.894	—	0.550
1.7	0.831	1.07	0.728
1.6	0.791	1.05	0.846
1.5	0.687	1.14	1.13
1.4	—	—	—
1.3	0.543	—	1.54
1.2	0.472	1.15	1.77
1.1	0.379	1.24	2.10
1.0	0.304	1.26	2.45
0.9	—	—	—
0.8	0.178	—	3.18
0.7	0.120	1.48	3.68
0.6	0.0878	1.36	4.07
0.5	0.0803	1.46	4.54
0.4	0.0371	1.6	5.10
0.3	—	—	—
0.2	0.0147	—	6.21
0.1	0.00950	1.5	6.71
0	—	—	—
-0.1	0.00350	—	7.84
0.2	0.00212	1.7	8.40
0.3	—	—	—
0.4	0.000613	—	9.78
0.5	0.000904	2.0	10.54
0.6	0.000132	2.3	11.45
0.7	0.000583	2.3	12.34
0.8	0.0000241	2.4	13.29
0.9	0.0000053	4.6	14.91

In the experiments of Richardson and Brotherton a very interesting fact was observed, namely, the difference between the apparent contact potential, as determined from the properties of the chemically emitted electrons, and from the photoelectric measurements. In the greater number of experiments the difference was about 0.6 volt, and this fact is difficult to explain on the assumption of a Maxwell distribution of energy among the electrons. With a

Maxwell distribution the straight line representing the relationship between  $\sigma V$  and  $V_1$  should intersect the  $V_1$  axis at  $V = 0, \pm e$ , at  $V_1 = -K$ , as  $V = V_1 + K$ , where  $K$  is the contact difference of potential. As the straight line on figs 1 and 2 assumes, with the new interpretation, the significance of an accidental tangent there is no reason to expect the intercept on the  $V_1$  axis, between the slope line and the origin, to represent the contact potential difference.

The observed difference between the photoelectric and the chemical zero can thus be considered as a confirmation of the present conclusion that the distribution of energy is not Maxwellian.

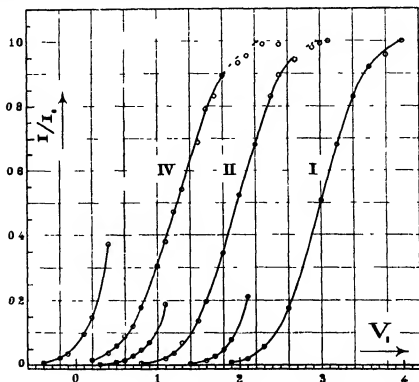


FIG 3

Fig 3 shows three characteristic curves  $i/i_0 = f(V_1)$  which have been selected because the measurements they represent were taken with the utmost care. In each case the smaller currents against the higher retarding potentials are shown separately to the left of the main curve. For these extensions of the main curve the ordinates have one-tenth of the value shown on the scale.

The curves correspond to the following pressures Curve I,  $p = 2.2 \times 10^{-5}$  mm., curve II,  $p = 1.4 \times 10^{-5}$  mm., curve IV,  $p = 5.0 \times 10^{-5}$  mm. Curve IV corresponds to the actual  $V_1$  axis, curves II and I are shifted to the right from IV by 1 and 2 volts respectively. It is important to note that in spite of the different pressures under which they have been obtained and the different surface conditions of the alloy resulting therefrom the curves have a very constant shape. The same thing is shown by the curves in Part I throughout the whole range of pressures from  $10^{-7}$  to  $10^{-3}$  mm. The slope of the middle part of the curves is almost a straight line. This fact distinguishes them very strongly from thermionic curves and shows a similarity with photo-electric ones.

Fig. 4 shows the corresponding differential curve  $\Delta(i/i_0) = \phi(V_1)$ , at  $\Delta V = 0.1$  volt. It represents the distribution of total energy among the

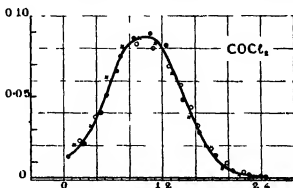


FIG. 4

emitted electrons. The points marked ● are derived from fig. 3, curve I, those marked × from fig. 3, curve II, and those marked ○ from fig. 3, curve IV. As the value of the contact potential difference is not known, we do not yet know the exact position of the zero on the  $V_1$  axis, since  $V = V_1 + K$  and when  $V = 0$ ,  $V_1 = -K$ . This point has been determined by experiments with mixtures of gases, which will be described in a later paper. These experiments show that the point of departure of the curves, such as those in fig. 3, from the saturation value  $i_0$  corresponds to the value  $V = 0$  within the limits of experimental error. The curves in fig. 3 have all been drawn to fit this experimentally determined zero. The experimental points show some systematic deviation from the curve, but it is doubtful if this exceeds the experimental error which is greatest in this part of the curve (see § 4, p. 62).

The curves in fig 3 give the following values for the contact potential difference and for the maximum electron energy  $E_m$  —

Curve I	$K = -2.0 \pm 0.1$ volts, $E_m = 2.5 \pm 0.1$ volts
Curve II	$K = -1.9_6 \pm 0.1$ volts, $E_m = 2.5 \pm 0.1$ volts.
Curve IV	$K = -2.2_4 \pm 0.1$ volts, $E_m = 2.6 \pm 0.1$ volts

Average value for  $E_m = 2.5_3 \pm 0.1$  volts

If  $N(V)dV$  denotes the proportion of the whole of the emitted electrons, which have energies within the range  $eV$  to  $e(V+dV)$ , the distribution function  $N(V)$  for  $\text{COCl}_2$  satisfies the equation

$$N(V) = Ae^{-k(V-V_m)^2}, \quad (5)$$

where  $A$  and  $k$  are constants, to within the degree of accuracy of the experimental data, or nearly so, except in the very high and very low energy parts of the spectrum. We can test this as follows, taking logarithms of (5) we have

$$\log_{10} A = \log N(V) + \frac{k}{2.30_3} (V - V_m)^2 \quad (6)$$

Thus the right-hand side of (6) should be a constant quantity for all values of  $V$ . When the data of fig 4, for example, are subjected to this test the result is Table II. In this table the value of  $k$ ,  $2.303 \times 1.087$ , has been determined

Table II — Test of the equation  $N(V) = Ae^{-k(V-V_m)^2}$  for  $\text{COCl}_2$

$V$ (volts)	0.2	0.4	0.6	0.8	1.0	1.2	1.4	1.6	1.8	2.0
$(V - V_m)^2$	0.64	0.36	0.16	0.04	0	0.04	0.16	0.36	0.64	1.0
$\log_{10} N(V)$	2.301	2.562	2.796	2.917	2.937	2.906	2.771	2.525	2.176	1.756
$\frac{k}{2.30_3} (V - V_m)^2$	0.696	0.391	0.174	0.044	0	0.044	0.174	0.391	0.696	1.087
$\log_{10} A$	2.997	2.953	2.970	2.961	2.937	2.950	2.945	2.910	2.872	2.843

by subtracting the values of  $\log_{10} N(V)$  at  $V - V_m = 0$  and  $V - V_m = 1$ , and thus eliminating  $A$ . It will be seen that the values of  $\log_{10} A$  in the last line of Table II, which are the sums of the quantities in the two preceding lines in the same column, vary very little over a range of  $V$  of 1.8 volts. The mean of these values of  $\log_{10} A$  is 2.924. It is probable that a more accurate value of this empirical constant will be given by the value of  $\log_{10} N(V)$  at which it is a maximum. This is 2.937. These two values do not differ much. It is possible that the average of them, 2.930, will be more correct.

than either. If we use this average value to re-determine  $k$  in combination with the data for  $V = 1.0$  and  $V = 2.0$  volts we find

$$k = 2.30_2 \times 1.15_2 = 2.65_2 \text{ volt}^{-2}$$

Equations (5) and (6) are only to be regarded as *empirical* equations which express the experimental data with a fair degree of accuracy. In all probability they do not represent the correct mathematical form of the distribution function. Nevertheless, they have a certain degree of practical utility. It will be noticed that equation (5) makes the distribution function completely symmetrical about the line  $V = V_m$ . That the actual distribution function for  $\text{COCl}_2$  is very nearly symmetrical about this line is one of its most striking features.

The extent of the deviation of the actual distribution functions from a Maxwell distribution is shown in fig. 8. In this figure the circles represent the experimentally determined values whilst the crosses give the values of the Maxwell distribution function  $N(V) = 0.775 V^{2e-2.2V}$  which fits the experimental data from  $V = 0$  to a little beyond  $V = V_m$ . It is at once seen that after passing the maximum the actual electron density becomes smaller and very rapidly enormously smaller than that which would be given by a Maxwell distribution formula.

### § 3 *Influence of the Geometry of the Electrodes on the Experimental Energy Distribution Curves*

The experiments described so far were all made with small drops of the alloy at the centre of a platinum cylinder 4.2 cm. long and 2.0 cm. in diameter, as described in Part I. Inasmuch as we are only concerned with the distribution of total kinetic energy the shape of the surrounding negative electrode is of no account provided it completely surrounds the small source. For this particular case the results are the same as for a small source at the centre of a charged sphere, where the geometry permits an easy calculation of the details of the motion of the electrons. In the actual experiment it is not possible to use a surrounding electrode with a completely closed surface as it is necessary to have openings for the tube to admit the alloy and for the escape of the drops. The presence of these apertures introduces an element of uncertainty into the analysis of the data which it is necessary to remove.

For concentric spherical electrodes it is possible to estimate the errors introduced by the finite size of the central electrode and by the apertures in the outer electrode. The problem of the application of spherical condensers

to energy distribution analysis has, in fact, been considered in detail by Lukirsky\*. The error introduced by the finite radius of the inner sphere is equal to the square of the ratio of its radius to that of the outer sphere. Thus in the experiments of Richardson and Brotherton (*loc cit*), where a sphere of 3.8 cm diameter was used and the drops had a radius of 0.3 to 0.4 cm, the error from this source would not exceed 1%. For concentric spheres, since all the trajectories are radial, the proportion of the electrons which escape through the apertures is simply the proportion which the solid angle subtended at the centre by the apertures bears to  $4\pi$ . With Richardson and Brotherton's electrode this would amount to about 2%. This error will be somewhat affected by the distortion of the radial field caused by the presence of the apertures, but it is not believed that this effect will be of much importance in the present experiments.

With electrodes of other shapes it is much more difficult to estimate the errors which may be introduced by the presence of apertures. We have considered two problems,† (1) a narrow emitting plane strip of indefinite length lying in the middle of a coplanar conducting strip of the same indefinite length and width  $2l$ , opposite to an opposing parallel plane electrode of the same dimensions,  $2l \times \infty$ , at a distance  $b$ , and (2) a small emitting source treated as a portion of a cylinder of radius  $a$  placed at the centre of a co-axial cylinder of radius  $b$  and length  $2l$ . In each the emitting electrons are endowed with a Maxwell distribution of kinetic energy.

It might be thought that problem (1) would be very simple. It is easy enough to obtain an integral, which expresses the number of electrons which escape through the aperture, but it is not integrable in finite terms, and we have not been able to find any series expansion for it which is of any practical use. However, it has one simple property. Every element of the integral is real and positive. By the artifice of multiplying the integrand by a certain factor which is positive and greater than unity throughout the domain of integration we can reduce it to a very simple integral which must thus have a value greater than that of the first integral. In this way we can show that the proportion of the emitted electrons which escape through the apertures between the planes must be less than

$$\frac{2}{\sqrt{\pi}} \sqrt{\frac{Ve}{kT}} \exp\left(-4 \frac{l^2}{b^2} \frac{Ve}{kT}\right) \quad (7)$$

\* 'Z. Physik,' vol. 22, p. 351 (1924).

† The details of these calculations may be published elsewhere.

except when  $Ve/kT$  is small, let us say appreciably less than unity. In (7)  $V$  is the opposing potential difference,  $e$  the electronic charge and  $k$  Boltzmann's constant. For any useful values of  $l/b$ , such as would correspond to the dimensions of apertures used in actual experimental arrangements, (7) is a very small quantity. It is zero when  $Ve/kT = 0$ , has a single maximum value  $\frac{1}{2\pi e} \frac{b}{l}$  when  $\frac{Ve}{kT} = \frac{b^2}{8l^2}$  and approaches zero with great rapidity as  $Ve/kT$  increases in the direction of  $+\infty$ .

In problem (2) it is possible to write an integral which expresses the number of electrons which escape from the ends of the cylinder. But the integral involves the smallest real root of a transcendental equation which so far has resisted all efforts to contract an infinite series which forms part of it. However, by a graphical method, it is possible to show that with the dimensions of the cylinder used in the present experiment the error in the experimental distribution function caused by the electrons which escape from the aperture is always very small for values of  $V$  at which the deviation from the Maxwell distribution, as shown, for example, in fig. 8, becomes appreciable.

The foregoing considerations make it highly improbable that the deviations from the Maxwellian distribution at high energies can arise from the effects of apertures in the surrounding electrode or from the finite size of the source. However, it is not possible to make calculations for the precise structure of electrodes and resulting distribution of electric field used in the present experiments so that the reality of the deviations cannot be regarded as completely established by these arguments.

We can obtain some further evidence bearing on this question by considering some of the older experiments. In 1921 it was proved by one of us\* that, within the limits of experimental error, the substitution for a spherical electrode of a cylindrical one of approximately the same form as the cylinder used in the present experiments did not alter the form of the characteristic curves. Again, if we compare the results obtained by Richardson and Brotherton (*loc. cit.*) when using a spherical electrode with those got with the cylinder in Part I and in the present experiments there is no appreciable difference in the range in which they overlap. However, this evidence is also not entirely conclusive because in the 1921 experiments the accuracy of measurement was far below what it is now, and in the experiments of Richardson and Brotherton the amount of overlap in the high energy region where the deviations occur is very slight.

\* Richardson 'Phil. Trans.,' A, vol. 222, p. 1 (1921)



#### § 4 Experiments with a Spherical Electrode.

In order to remove any uncertainty, resulting from the considerations set out in § 3, as to the reality of the deviations from the linear  $\sigma V$   $V_1$  plots, we decided to repeat the experiments using a spherical platinum electrode, with the same refinements and precision as had been attained with the cylinder, and extending well into the high energy region. With a spherical electrode, as we have pointed out, the disturbing effects of apertures and finite size of drops can be ascertained with sufficient certainty and shown to be unimportant.

The electrode used was a seamless pure platinum sphere about 2.75 cm. in

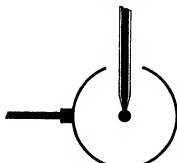


FIG 5

diameter with two circular apertures each 1.00 cm. in diameter. Fig. 5 represents exactly the actual size of the spherical condenser. It also indicates approximately the maximum size of the drops. This depends somewhat on the time  $T$  of a drop. The first experiment was performed with  $T = 2$  minutes 30 seconds and the maximum diameter  $d$  was about 0.4 cm. In other experiments  $T$  was about 10 seconds and  $d$  about 0.3 cm. With this value of  $d$  the ratio between the diameters

of the sphere and the drop is  $27.5/3 = 9.2$  and the error introduced by the finite size of the source is thus about 1%. The combined error arising from this and from the apertures was in the neighbourhood of 3–5%.

The sphere was thoroughly cleaned with benzene, caustic soda, nitric acid, and distilled water, it was then placed in a quartz tube and evacuated for many hours at  $1000^\circ \text{C}$ . In the actual experiment the spherical electrode was used without a heating arrangement and for that reason the pressure of the active gas was not allowed to exceed  $5 \times 10^{-5}$  mm. At such low pressures the behaviour of the outer electrode, as regards variations of the contact potential difference, was the same as that with the cylindrical electrode with a heating arrangement. This is important because changes in the contact potential difference affect the experimental determination of the characteristic curves and it is necessary to understand and allow for these effects.

As a result of a large number of observations on a variety of reactive gases we can sum up the effects of these gases on the contact potential difference between the alloy and the platinum electrode as follows:

At very low pressures the variation of the contact potential difference is determined mainly by adsorption of the active gas on the platinum surface. This effect is quite reversible. The active gas is electronegative and it reduces, when adsorbed, the work function of platinum, so that the contact potential difference is less than in a good vacuum. At the lowest pressures, of the order  $10^{-6}$  mm, the equilibrium conditions result in the formation of a monomolecular layer on the platinum surface and the effect of the gas in reducing the work function of the platinum is then the greatest. With increase of pressure the equilibrium conditions allow the metal surface to be covered with a layer more than 1 molecule thick and this will reduce the effect of the first layer of molecules. The contact potential difference will thus show an increase with pressure (maximum about 0.5 volt, see Part I, figs 3 and 4, pp 36-37). When the pressure of the active gas exceeds  $5 \times 10^{-3}$  mm, the vigour of the chemical reaction with the alloy is increased so much that it may cause individual molecules of the alloy, or the reaction products, to evaporate and condense on the platinum surface. This is shown by the fact that, even if the exposure to the gas at these higher pressures lasts only a short time, a thin layer can be noticed on the inner side of the platinum electrode after the apparatus has been taken to pieces. In this respect, apparently, water vapour gives the greatest effect. When such layers are present complicated permanent, or semi-permanent, changes in the contact potential difference may occur of the type described by Richardson and Brotherton (*loc cit*, pp 39-41) as hysteretic effects. It is probable that an essential feature of the mechanism of these effects is the presence on the platinum of a layer of reaction products which insulates it from the electrons and thus forms a condenser of very high capacity.

Three independent experiments were carried out with the spherical electrode, the  $\sigma V$   $V_1$  plots of which are given in fig 6. Experiment 1 (marked 2 in fig 6). Time of drop  $T = 2$  minutes 30 seconds, time of exposure  $dt = 60$  seconds, pressure  $p \sim 2.8 \times 10^{-5}$  mm, saturation current  $i_0 = 3.7 \times 10^{-10}$  amp. Experiment 2 (1 in fig 6).  $T = 7-8$  seconds,  $dt = 30$  seconds,  $p \sim 1.7 \times 10^{-5}$  mm,  $i_0 = 3.8 \times 10^{-9}$  amp. Experiment 3 (3 in fig 6).  $T = 9.5$  seconds,  $dt = 35$  seconds,  $p \sim 4.7 \times 10^{-5}$  mm,  $i_0 = 9.2 \times 10^{-9}$  amp. Curves 2 and 3 in fig 6 are shifted to the right relative to curve 1 by 0.2 and 1.2 volts respectively. It will be seen that the results of all three experiments are consistent with each other and, within the limits of experimental error (maximum 5%), the plots can be very well superimposed on each other and on those obtained with the cylindrical electrode. This is shown on curve 2 of

fig 6 where the crosses represent the experimental points of curve II, fig 1, which was obtained with the cylinder

*This proves that the apparatus with the cylindrical electrode does give the correct energy distribution in the spectrum of the emitted electrons*

As the cylinder is more conveniently adapted to heating and we thus avoid troublesome changes in the contact potential difference, we have used it in preference to the sphere in most of the other experiments

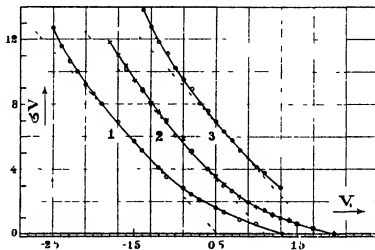


FIG 6

The actual characteristic curves corresponding to 1 and 2 in fig 6 are shown as 1 and 2 respectively in fig 7 where curve 2 is shifted to the right relative to curve 1 by 1.0 volt. In each the high energy part is shown separately from the rest with the vertical scale magnified ten times. The irregularity of the points in the neighbourhood of  $v/v_0 = 1$ , which is particularly noticeable in curve 1, fig 7, is due to small changes in the contact electromotive force which occur during the experiment, coupled with the fact that the absolute error of measurements is greatest in this part of the curve. It is impossible to determine the exact course of this part of the curve with anything like the certainty which is attained in the high energy part, on the left-hand side.

Figs. 8 and 9 show the energy distribution curves  $\Delta(v/v_0)/\Delta V = N(V)$  for  $\Delta V = 0.1$  volt corresponding to curves 1 and 2 respectively of figs 6 and 7. The points, which are measured from fig 7, are shown as circles. In fig 8 the crosses represent the Maxwell function  $0.775 V^2 e^{-2.4V}$  which fits the low energy part of the curve. The deviation of the high energy part from this function is very striking. The crosses in fig 9 are the corresponding data got

in one of the experiments with the cylindrical electrode, and shown as solid dots in fig 4

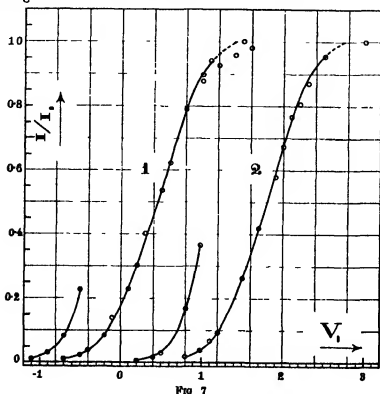


FIG 7

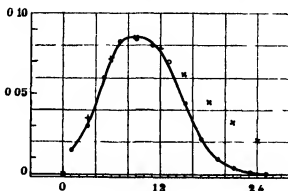


FIG 8

Particular attention was paid to ascertaining whether small accelerating electric fields have any real effect in increasing the electron emission. It will be remembered that the measurements with the cylindrical anode in

Part I gave a negative answer to that, but there were some elements of uncertainty arising from the geometry of the cylindrical electrode. It will be seen from Table III that the electron current (in arbitrary units) remains constant within 3% with accelerating fields up to +6 volts. The current was saturated at about +1.8 volts. These results were obtained in the course of the experiment 1 of figs. 6, 7, and 9.

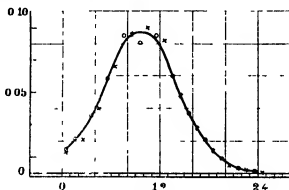


FIG. 9

Table III—Test of Effect of Accelerating Field on Saturation Current

Volts $V_1$	+2.0	+3.0	+4.0	+5.0	+6.0
Current $I$	195	194	199	195	196

### §5 *The Low Energy Part of the Spectrum, the Determination of the True Zero on the Volt Scale and of the Maximum Energy*

In this section we outline very briefly the conclusions we have reached on these questions. It is impossible at this stage to give the reasons for them adequately as they depend very largely on an accumulation of evidence which has been got from the investigation of gases other than  $\text{COCl}_2$  and which will be described in the papers to follow.

At the end of Part I we expressed a suspicion that the deviations from linearity of the  $\sigma V - V_1$  plots in the low energy part of the spectrum, i.e., near  $V = 0$ , were a real feature of the energy distribution and not due to secondary complicating causes tending to prevent the attainment of saturation, as had hitherto been supposed. We are now quite convinced that these deviations are real. They are found to persist unchanged, as we mentioned in Part I, over an enormous range of variation of different experimental con-

ditions. The former straight lines are now only tangents to a curve which is continuous from  $V = 0$  to the highest values of  $V$  for which measurements can be made, but which happens to be nearly a straight line over a considerable range. The curious conflict found by Richardson and Brotherton (*loc cit*) between the voltage zero when determined photoelectrically and when determined from the  $\sigma V$   $V_1$  plots on the old interpretation now disappears. In fact, we now know where the true voltage zero is. It coincides with the photoelectrically determined zero and with the point at which the current begins to drop from the saturation value. In this respect we are back at the position reached in 1921.\* The  $\sigma V$  plots have to reach this value of  $V_1$  at  $\sigma V = 0$  and they approach it by means of the continuous curve found experimentally and not by means of a straight line which leads somewhere else. Other facts which make it quite certain that the deviations of the linear plots near  $V = 0$  are real are derived from experiments with gases other than  $\text{COCl}_2$ . The shapes of these "feet" are found to be characteristic of the different gases, in some gases they are even absent and the  $\sigma V$  plots are linear down to  $V = 0$ , to within the limits set by the errors of measurement.

We therefore conclude that the diagrams such as figs. 4, 8, and 9 represent the true distribution of energy among the emitted electrons, to the limits of accuracy set by the experimental measurements.

The reason we are so sure that the true voltage zero coincides with that determined photoelectrically and with the voltage at which the current begins to fall from the saturation value is because we have devised a new experimental method of attacking this problem which confirms those determinations. This method depends on experiments with mixtures of different gases and will be described in a later paper. The reason why this position was abandoned previously is because in the older experiments, owing to the difficulties created by changes in the contact potential difference, the point of departure from saturation could not be determined with sufficient accuracy to enable a decision to be made as to which of the two zeros then measured experimentally was the correct one.

Although we can state quite definitely, as we have done above, that the deviations from linearity of the  $\sigma V$   $V_1$  plots in the low energy part of the spectrum are a real feature of the energy distribution and not due to secondary complicating causes tending to prevent the attainment of saturation, it is necessary to point out that sometimes such secondary factors preventing the attainment of saturation did exist in the immediate neighbourhood of the

\* Richardson, *loc cit*

**actual zero** These distortions of the extreme low energy part of the electron spectrum were observed only at pressures higher than  $10^{-5}$  mm, and it seems that the higher the pressure the more readily they appear. Curve 1 of fig 7, obtained at a pressure of  $1.7 \times 10^{-5}$  mm, gives a good idea of the magnitude of these distortions. The effect can be explained by a greater change in contact potential difference at higher pressures during the experiment. By all indications, such distortions existed in some greater magnitude in the higher pressure experiments of Richardson and Brotherton (*cf* fig 1, p 25, of their paper). They have probably played an important role in making it much more difficult to give a correct interpretation of the difference between the chemical and photoelectric zeros.

An analysis of the energy distribution curves, such as I, II, and IV of fig 4, and the curves of figs 8 and 9, shows that the curves do not plunge into the volt axis very sharply in the neighbourhood of the maximum energy, not nearly so sharply as they do in the case of the photoelectric effect. However, the detailed investigation of the high energy part of the electron spectrum (see § 2) has proved quite definitely that the distribution curves do *not* approach the volt axis *asymptotically*. This can be very well seen, for instance, from curve III of fig 1 and curve IV of fig 2. With increase of the energy  $V$  the tangent of the curves approaches a value equal to infinity. It follows from this that the *actual* maximum energy  $E_a$  can be estimated from the  $\sigma V$   $V_1$  plots and could be defined as the value of  $V_1$  corresponding to  $\sigma = \infty$ . There is some reason, however, for dealing with the "*effective*" maximum energy rather than with the actual maximum energy  $E_a$ . This effective maximum energy  $E_m$  will be defined as the value of  $V_1$  at which the energy distribution curve *practically* touches the volt axis. Below we give the values of  $E_m$ , obtained from the energy distribution curves, and the corresponding values of  $v/v_0$ .

Fig 4, curve I	$E_m = 2.5$ volts, $v/v_0 = 0.0008$
Fig 4, curve II	$E_m = 2.5$ volts, $v/v_0 = 0.0006$
Fig 4, curve IV	$E_m = 2.6$ volts, $v/v_0 = 0.0006$
Fig 8, curve 1	$E_m = 2.5$ volts, $v/v_0 = 0.0006$
Fig 9, curve 2	$E_m = 2.5$ volts, $v/v_0 = 0.0005$
Average value for $E_m = 2.5$ volts	

The values of  $E_a$ , obtained from the  $\sigma V$   $V_1$  plots are —

Fig 1, curve III	$E_a = 3.0$ volts, $v/v_0 < 0.000005$
Fig 2, curve IV	$E_a = 3.1$ volts, $v/v_0 < 0.000005$
Average value for $E_a = 3.0_5$ volts	

It will be seen from this that the values of  $E_m$  are about 0.5–0.6 volt higher than  $E_m$ , but the electron current at  $V = E_m$  is already less than 1/1000 of its saturation value and, as can be seen from the  $\sigma V - V_1$  plots, it falls down very quickly at  $V > E_m$ , so that there is every reason to treat the part of the distribution curve at  $V > E_m$  as a "tail" caused by some secondary factors. With many other gases it was even impossible to obtain the ratios  $i/i_0$  at  $V > E_m$  because of the small absolute value of  $i_0$ , and thus it was impossible to find the actual maximum energy  $E_m$ .

It may be pointed out that the values of  $E_m$  can be approximately estimated also from  $\sigma V - V_1$  plots. As can be seen from figs 1, 2, and 6, at  $V = E_m$  the curves turn sharply to the right from the tangent indicated in the diagrams. According to this method the curves III and V of fig 1 (these curves were obtained in the investigation of the extreme high energy part of the electron spectrum and because of the lack of the middle points it was impossible to plot differential curves for those data) give the following values of  $E_m$  —

Fig 1, curve III	$E_m = 2.4$ volts, $i/i_0 = 0.0010$
Fig 1, curve V	$E_m = 2.5$ volts, $i/i_0 = 0.0007$

### § 6 The Relation between the Maximum Energy of the Emitted Electrons and the Energy of the Reaction.

In the reaction between  $\text{COCl}_2$  gas and  $\text{K}_2\text{Na}$  alloy we are concerned with the formation of a polar chemical bond between the Cl atom, which on the ground of our experimental results we can assume to be simultaneously liberated from the rest of the molecule and converted into a negative ion, and one of the positive K, or Na, ions of the metal, which is chemically bound to other atoms of the metal. It is important to realize that the energy of the formation of the bond will be dissipated, among a number of the adjacent metal atoms, in a very short time, of the order of  $10^{-13}$  seconds, because of the normal heat conductivity of the metal\*. On the other hand, the time required for an adiabatic formation of a bond is of the order of at least  $10^{-13}$  seconds. Therefore, the energy developing during an adiabatic formation of the bond will be practically wholly dissipated to the adjacent metal atoms by the time the incident molecule first reaches the equilibrium position corresponding to the maximum negative potential energy of the nascent bond. We thus arrive at an important conclusion that *the observed chemical electron emission can only*

\* Langmuir, 'Phys. Rev.', vol. 8, p. 149 (1916), 'J. Amer. Chem. Soc.', vol. 38, p. 2233, (1916)



*be a result of a spontaneous rearrangement of the electronic system of the nascent bond*

The experimental evidence on the spectra of alkali-halide molecules,\* as well as the results of theoretical considerations of non-adiabatic chemical processes,† lead to the conclusion that the formation of a stable alkali-halide molecule in the normal state from the two neutral atoms is an essentially non-adiabatic process. That is, if we consider the reverse process of dissociation it is necessary for the alkali-halide molecule in the ground state to absorb a light quantum to give dissociation into normal atoms. It is pointed out, however, that the chemical transformation may be concerned with a quantum transition which involves no change of the character of the electronic term, since in the case of alkali-halides the ground state (which gives dissociation into ions) and the first excited state (which gives dissociation into normal atoms) may both be  $^1\Sigma$ -terms. If we accept that, in our case of the heterogeneous formation of the polar bond, the process of formation is, in principle, analogous to that in the case of the alkali-halogen molecules, it will follow that the chemical electron emission has as its basis a pure quantum effect.

As the amount of energy liberated in most chemical reactions between a gas and a metal is of the order of a few volts, it will be safe to conclude that no chemical electron emission from a metal can be observed during an adiabatic heterogeneous chemical transformation, which does not involve a spontaneous rearrangement of the atomic electrons, such as, for example, the condensation of metal atoms on the surface of their own metal.

So far as we are only concerned in giving an energy account of the chemical process it is not necessary to know exactly the actual mechanism of the formation of the polar bond, but we have to know the initial and final states of the system. Because of the short time of de-excitation of the polar bond formed the final state of the transformation will be that which corresponds to the instant of formation of the bond. It is evident that a direct comparison of the maximum energy of the emitted electrons with the chemical energy calculated merely on the basis of thermodynamical arguments under equilibrium conditions will be wrong.

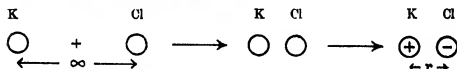
The exact theoretical calculation of the energy of the heterogeneous chemical transformation, corresponding to the instant of the formation of the polar bond, even in our comparatively simple case would be quite hopeless. It

\* Sponer, "Molekülstruktur," p 105, Leipziger Vorträge (1931)

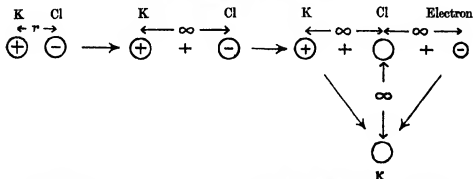
† London, 'Z Physik,' vol 46, p 455 (1928), and vol 74, p 143 (1932)

would be necessary to determine the distance between the ions of the polar bond at the instant of its formation and to calculate the effect of the adjacent metal atoms upon the bond considered. Neither of these two factors can be estimated accurately. We should like, therefore, to put forward an approximate method which will enable us to estimate the order of magnitude of the amount of energy liberated at the chemical transformation. One great simplification which can be introduced in the heterogeneous change considered is that the problem of interaction between alkali and halogen atoms may be simply reduced to an electrostatic problem of interaction between point charges. In fact, the ions constituting the chemical bond have electronic systems like inert gases, and it is easy to show that at the interatomic distances involved, the deformation (polarization) of the ions of the bond by each other, or by the positive ions and free electrons of the metal, will be small. Therefore, their electric fields will be those due to point charges  $\pm e$  at the centre of each.

Consider first the simpler case of the combination of a neutral atom of K, for example, with a neutral atom, let us say, of Cl at the instant when the electron jumps from the K to the Cl atom at the distance  $r$  between their centres. We can represent this process by—



Let us denote the overall energy liberated in this process by  $(E_s)$ . Now consider the following reverse process, by an alternative route—



The dissociation of  $\text{K}^+\text{Cl}^-_{(r)}$  into  $\text{K}^+$  and  $\text{Cl}^-$  requires energy  $e^2/r$ , the neutralization of  $\text{Cl}^-$  requires energy  $A_{\text{Cl}}$  (the electron affinity of Cl) and the neutralization of  $\text{K}^+$  supplies energy  $I_{\text{K}}$  (the ionization potential of K). It follows that

$$(E_s) = e^2/r + A_{\text{Cl}} - I_{\text{K}} \quad (8)$$

or, in words, the energy liberated instantaneously in the reaction is equal to the mutual potential energy of the two ions at the instantaneous distance  $r$  plus the electron affinity of the negatively charged atom minus the initial negative energy of the displaced electron. This result is exact and will apply to any reaction of this type (except that, in general, the expression for the potential energy of the ions will be more complicated than  $-e^2/r$ ).

Now consider in what ways the actual heterogeneous reaction will differ from this homogeneous reaction. In the first place there will be a certain distance  $r_1$  characteristic for the heterogeneous reaction. We shall assume a normal collision diameter and put  $r_1$  equal to the sum of the radii,  $r_{Cl} + r_K$ , of the neutral halogen atom and of the "neutral atom" of the metal. For the latter quantity we shall take one-half of the closest distance between atoms in the metal. For the  $Cl^- - K^+$  bond we may take  $r_{Cl} = 1.07 \times 10^{-8}$  cm.\* and for  $r_K = 2.25 \times 10^{-8}$  cm,\* then  $r_1 = r_{Cl} + r_K = 3.32 \times 10^{-8}$  cm. This distance is probably larger than the equilibrium distance of a normal KCl molecule by about  $0.5 \times 10^{-8}$  cm. In the second place the displaced electron is no longer in the ground state of the K atom, it is in one of the states occupied by the free electrons in the metal. Adopting Sommerfeld's theory the minimum negative energy of an electron (at  $0^\circ$  K.) will be  $\phi = W_s - \mu$  (where  $\phi$  is the work function of the metal,  $W_s$  is the total height of the potential barrier and  $\mu$  is the energy associated with the highest possible state occupied by the free electrons) and this will have to replace  $I_K$  in equation (8). There is, however, another difference. According to fig. 10, which gives a schematic representation of the chemical bond at the instant of formation, an essential difference between this and that of a free molecule is that the negative Cl ion is subject not only to the attraction of the positive K ion, but also to that of the whole mass of metal to the right-hand side of the plane  $x = x_0$ . Since the negative charge is, in effect, in the plane  $x = x_0 + r_{Cl}$ , and if we make the further assumption that the field of force to the left of the plane  $x = x_0$  is determined by the mirror image of the charge, the final negative energy of the pair of ions will be increased by the amount  $\frac{e^2}{4} \left( \frac{1}{x_0 + r_{Cl}} \right)$ , where the parameter  $x_0$  is determined by the relation  $\phi = \frac{1}{4} \frac{e^2}{x_0}$ . Thus we obtain finally

$$E_s = \frac{e^2}{r_1} + \frac{e^2}{4} \left( \frac{1}{x_0 + r_{Cl}} \right) + A_{Cl} - \phi \quad (9)$$

\* Landolt-Bornstein, 'Physikalisch-Chemische Tabellen,' 1924-1931

This formula will only be approximately true, as the distortion of the electron distribution in the metal by the  $K^+$  ion which combines with the  $Cl^-$  ion will come into play at these small distances. This would tend to make the expression  $\frac{e^3}{4} \left( \frac{1}{x_0 + r_{cl}} \right)$  too large. However, there are other uncertain elements which come in at these distances, arising from our ignorance of the precise form of the electron potential energy distribution as we cross the surface of the solid, and it is probable that, in fact, equation (9) is very little in error. In particular, it is fairly certain that it is less incorrect than the combined inaccuracies of the data to which it is applied.

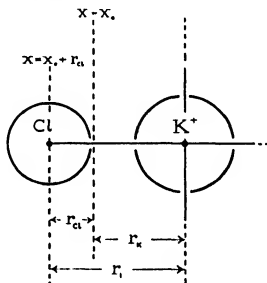


FIG 10

Since the actual initial state is the gaseous  $(COCl_2)$  molecule, and not the free (Cl) atom, it is necessary to take into account the energy of the corresponding dissociation of  $(COCl_2)$ . Dissociation of the gaseous molecule will occur simultaneously with the electronic rearrangement of the nascent polar bond, on account of the weakening of the bond connecting the Cl atom to the rest of the  $COCl_2$  molecule which is caused by the electron transference. It is well known\* that in the photochemical decomposition of molecules both the dissociation energy  $D$  and the energy of electron transference  $\epsilon$  have to be included in the photochemical equation  $h\nu = \epsilon - D$ , thus showing that

\* Cf Franck, 'Naturwiss.', vol. 10, p. 217 (1931).

dissociation occurs *simultaneously* with the electronic rearrangement. We have been able to establish, from a comparative study of a large number of different reactions, that in the case of  $(\text{COCl}_2)$  the energy of dissociation  $D$  of the process  $(\text{COCl}_2) = (\text{CO}) + 2(\text{Cl}) - D$  is equally distributed between the two Cl atoms. This shows that two Cl atoms of the  $(\text{COCl}_2)$  molecule normally enter into the reaction with the metal simultaneously.

Relation (9) will then take the form

$$E_e = \frac{e^2}{r_1} + A_{\text{Cl}} + \frac{e^2}{4} \left( \frac{1}{x_0 + r_{\text{Cl}}} \right) - \phi - \frac{1}{2}D \quad (10)$$

The electron affinity  $A_{\text{Cl}}$  of the (Cl) atom can be derived from the grating energy of the alkali halogen crystals, it is  $A_{\text{Cl}} = 3.8 \pm 0.3$  volts\* (A more accurate determination of the electron affinity has been made only for the iodine atom, by the method based on the thermal equilibrium at high temperature of CsI vapour).

It may be pointed out that the energy of the reaction  $E_e$  can be put in a direct connection with the thermochemical data if we make use of the relation between the grating energy  $U_{\text{KCl}}$  and the electron affinity  $A_{\text{Cl}}$ . This relation is†

$$U_{\text{KCl}} = Q_{\text{KCl}} + \frac{1}{2}D' + I_{\text{K}} + S_{\text{K}} \quad A_{\text{Cl}},$$

where  $Q_{\text{KCl}}$  is the heat of the reaction  $\frac{1}{2}(\text{Cl}_2) + [\text{K}] = [\text{KCl}]$  at  $0^\circ \text{K}$ ,  $D'$  is the heat of dissociation of  $\text{Cl}_2$ ,  $I_{\text{K}}$  is the ionization potential of the (K) atom,  $S_{\text{K}}$  is the heat of sublimation of [K] at  $0^\circ \text{K}$ . Now,

$$U_{\text{KCl}} = M_{\text{KCl}} + S_{\text{KCl}} \approx \frac{e^2}{r_0} + S_{\text{KCl}},$$

where  $M_{\text{KCl}}$  is the energy of the process  $(\text{Cl}^-) + (\text{K}^+) = (\text{KCl})$ , which is approximately equal to  $e^2/r_0$ ,  $r_0$  being the equilibrium interatomic distance of the (KCl) molecule in the normal state, and  $S_{\text{KCl}}$  is the heat of sublimation of [KCl] at  $0^\circ \text{K}$ . We thus obtain

$$E_e = (Q_{\text{KCl}} + \frac{1}{2}D') + (I_{\text{K}} - \phi) - (S_{\text{KCl}} - S_{\text{K}}) + \left\{ \frac{e^2}{x_0 + r_{\text{Cl}}} \right\} - e^2 \left( \frac{1}{r_0} - \frac{1}{r_1} \right) \quad (11)$$

Taking for  $\frac{1}{2}D = 1.83$  volts, and for the work function of the  $\text{K}_2\text{Na}$  alloy  $\phi_{\text{K}_2\text{Na}} = 2.5 \pm 0.1$  volts (Richardson, *loc. cit.*), and therefore  $x_0 = 1.44 \times 10^{-8}$  cm, we obtain, according to relation (10),

$$E_e = 4.34 + 3.8 + 1.44 - 2.5 - 1.83 = 5.26 \text{ volts}$$

\* van Arkel and de Boer, "Chemische Bindung als elektrostatische Erscheinung," p. 63 (Leipzig, 1931).

† *Ibid.*, p. 56.

with uncertainty at least  $\pm 0.3$  volt. The calculation according to relation (11) gives practically the same value for  $E_e$ , it is less only by 0.1 volt.

We obtain the maximum possible value for the energy of the emitted electrons if we consider the de-excitation of the nascent chemical bond by collision of the second kind with an electron of maximum energy  $\mu$  inside the metal (at 0° K). This will be equal to

$$(E_e - \phi_{K,Na})_{max} = 2.7,$$

It may at once be seen that the calculated value of the maximum energy of the emitted electrons is of the same order of magnitude as the observed, which lies between 2.5<sub>e</sub> and 3.0<sub>e</sub> volts.

Since the potential energy of the bond  $Cl^- - K^+$  is in inverse proportion to the distance  $r_1$  between the centres of the ions and therefore it is sufficient if the distance is decreased by about 10% ( $\Delta r_1 \approx 0.4 \times 10^{-8}$  cm) to give an increase of energy equal to 0.5 volt, it seems to be natural to assume that the observed unsharp limit for the maximum energy (the "tail") is a result of the uncertainty of the collision diameter. On the one hand, as gaseous molecules take part in the reaction the collision diameter will depend upon the orientation of the molecule at the instant of collision. It is hardly probable, however, that the fact that the  $COCl_2$  molecule contains two Cl atoms plays any important part, for the same unsharp upper limit of energies of the emitted electrons was observed, as we shall show in a later paper, also for (NOCl). On the other hand, the collision diameter will depend upon the electron atmosphere of the metal. Since the metal is at ordinary room temperature there exists a more or less sharp limit for the maximum kinetic energy of the free electrons in the metal, and we may expect that there will be a more or less abrupt outer limit for the electron atmosphere of the metal, at which most of the halogen molecules will undergo chemical transformation. However, there may be some halogen molecules which will penetrate this electron boundary and thus will undergo transformation at a shorter distance from the metal.

There are two more factors which will act in the same direction of giving an unsharp character for the distance  $r_1$ , and therefore for the upper limit of energies of the emitted electrons. One factor is the thermal agitation of the alloy atoms (ions), the  $K_2Na$  alloy used in the experiments is quite a mobile liquid at room temperature. Another is that the composition of the metal is not homogeneous. This latter fact may turn out to be rather important and not only because of the different size of the positive K and Na ions (the

radius of the  $\text{Na}^+$  ion is less by about  $0.3 \times 10^{-8}$  cm), but also because of the possible difference in the character of the interatomic forces of the two bonds  $\text{Cl}^- - \text{K}^+$  and  $\text{Cl}^- - \text{Na}^+$ .

We might also expect some part of the tail to be caused by the kinetic energy of the partners at the three-body collision, i.e., the gas molecule, the metal atom and the free electron. In particular, the estimation of this temperature effect of the free electrons, on the basis of the Fermi-Dirac distribution function for the electrons inside the metal at  $300^\circ \text{K}$ , shows that its contribution to the tail can be appreciable, but only at  $V - E_m < 0.2$  volt.

In concluding we should like to take this opportunity of acknowledging our indebtedness to the Department of Scientific and Industrial Research for a grant which has made it possible to carry out these investigations.

### *Summary*

This paper starts by summarizing some general conclusions we have reached from experiments on the reaction between  $\text{K}_2\text{Na}$  and 22 different gases. These include —

- (1) The energy distribution is not in general of the Maxwellian type.
- (2) The distribution curves all rise from a small value at zero energy ( $V = 0$ ) to a maximum at a certain energy ( $V = V_m$ ) and fall to zero at a certain maximum energy ( $V = E_s > V_m$ ).
- (3) These curves have a "tail," like that in the photoelectric effect, and  $E_s$  is approached rather gradually. A practical maximum energy  $E_m$  is obtained by disregarding the tails. About 99.9% of the electrons have energies below  $E_m$  ( $E_s > E_m > V_m$ ).
- (4) The distribution curves for energies  $> V_m$  and  $< E_m$  can all be represented quite closely by  $N(V)dV = A e^{-k(V-V_m)^c} dV$ ,  $A$ ,  $k$  and  $c$  constants. For the most energetic reactions this equation is a fair approximation from  $V = 0$  to  $V = E_m$ .
- (5) For the Cl compounds  $E_m + D = \text{constant}$ , where  $D$  is the dissociation energy of the relevant reaction.
- (6) At very low pressures the electron emission is proportional to the pressure. It usually rises to a sharp maximum at a quite low pressure and then falls.
- (7) The yield of electrons diminishes rapidly as the available chemical energy diminishes.

(8) The facts can be accounted for on the view that in the formation of the polar bond, which involves a spontaneous rearrangement of the electronic systems of the atoms concerned, the bond is stabilized spontaneously by a three body collision with a free metallic electron which carries away all the reaction energy. The effect is thus a quantum phenomenon. This view gives a reasonable account of the energy distribution. It also allows of a simple interpretation of the equation  $E_m + D = \text{constant}$ . It is  $E_m = E_s - \phi$ , where  $E_s$  is the energy of the corresponding elementary chemical reaction responsible for  $E_m$  and  $\phi$  is the work function of the metal. *This relation appears to be general and is similar to the fundamental law of photoelectricity.*

A more refined and detailed experimental investigation than has hitherto been carried out with phosgene is then given. This is followed by a discussion of the low energy part of the spectrum, the determination of the true zero on the volt scale and of  $E_m$  and its relation to  $E_s$ .

The last section confirms the equation  $E_m = E_s - \phi$  for the particular case of  $\text{COCl}_2$ .

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### *The Structure and Formula of 12-Phosphotungstic Acid.*

By J. F. KEGGIN, Ph.D.

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[PLATE I]

#### 1 Introduction

This acid belongs to a large class of compounds known as the heteropolyacids, of which the structures and exact formulæ have long been a subject for speculation. The heteropolyacids are compounds in which one atom of such elements as P, Si, As, B, Al, etc., is combined with a number of atoms of an element such as W or Mo, together with a relatively large number of atoms of oxygen. In addition to the elements mentioned, it has been shown that the oxides of a considerable number of other elements show a tendency to form heteropolyacids. All formulæ proposed indicate a relatively large number of atoms in the molecule, and a complex structure. In spite of the complexity of the molecule, many of these acids are quite stable, and form stable salts with practically all metals. The best known compounds of this group are the silicotungstic, silicomolybdic, phosphotungstic and phosphomolybdic acids, in which one atom of silicon or phosphorus is combined with a number of atoms



of tungsten or molybdenum. Mixed acids are also known in which a number of tungsten atoms are replaced by molybdenum atoms, or *vice versa*. The heteropolyacids are classified according to the ratio of the numbers of the two types of cations present.

Throughout the whole class of heteropolyacids, those acids which have the same cation-cation ratio tend to be isomorphous and have similar properties.

These acids have several outstanding properties, the following apply more especially to the 12-acids, though several of the other types of acids are very similar —

- (1) They form crystals containing large amounts of water of crystallization.
- (2) They are usually very soluble in solvents containing oxygen in the molecule\*—in water, ether, alcohols, alcoholic esters, ketones, aldehydes, etc., but not in such solvents as benzene which contain no combined oxygen. Ether will extract these acids from aqueous solution, forming a system of three layers, with ether on top, an aqueous layer next and at the bottom a saturated solution of the acid in ether. It has been suggested† that a loose compound of the oxonium type is formed with ether. This property is used in the preparation and purification of the heteropolyacids.
- (3) They are readily reduced‡ by common reducing agents such as nascent hydrogen,  $\text{H}_2\text{SO}_3$ , etc.
- (4) Perhaps the most remarkable property is the ability of these acids to form insoluble precipitates, often crystalline, with many substances, which substances often have large molecules. With the alkaloids bulky precipitates are formed, the standard method of estimating nicotine makes use of this property. They have been suggested as antidotes in cases of alkaloid poisoning. Albumen, peptones, etc., also give precipitates with these acids, which are used in their separation. They give precipitates with substances containing urea, and with amino-acids form crystalline compounds§. With basic dyes they give extremely insoluble lakes which are remarkably stable and fast to light.

The present work is concerned with 12-phosphotungstic acid.

\* Scroggie, 'J. Amer. Chem. Soc.', vol. 51, p. 1057 (1929).

† Rosenheim and Jaenicke, 'Z. anorg. Chem.', vol. 100, p. 319 (1917).

‡ Wu, 'J. Biol. Chem.', vol. 43, p. 189 (1920).

§ Drummond, 'Biochem. J.', vol. 12, p. 5 (1918).

## 2 Brief Review of the Literature with Reference to the Constitution of the 12-heteropolyacids

The heteropolyacids were first discovered by Berzelius\* in 1826. Among the more important of the earlier workers on these compounds, De Marignac,† Scheibler,‡ Sprenger,§ Soboleff,|| Kehrman,¶ Parmentier,\*\* and Copaux†† may be mentioned. Most of these workers suggest  $H_3PW_{12}O_{40}$  and  $H_4SiW_{12}O_{40}$  as the empirical formulæ of the anhydrous 12-phosphotungstic and 12-silicotungstic acids, as a result of dehydration experiments.

In 1908, Miolati and Pizzighelli,‡‡ and later Rosenheim and co workers,§§ suggested  $H_7[P(W_2O_7)_6] \cdot nH_2O$  and  $H_8[Si(W_2O_7)_6] \cdot nH_2O$  as the formulæ of these acids, as a result of titration experiments. This type of formula is known as the Miolati-Rosenheim formula, and is now considered as the classical formula.

In 1913 W and D Asch||| suggested a structure for the 12-acids, in which the molecule consists of two hexite rings. This gave  $H_3SiW_{12}O_{42} \cdot nH_2O$  as the formula for 12-silicotungstic acid.

Pauling,¶¶ in an attempt to explain the properties of the 12-heteropolyacids, proposed, on theoretical grounds, the first three-dimensional structural formula in which the positions of the atoms and the exact manner of co-ordination were indicated. After reviewing the literature, he decided that the 12-phospho-acids are three-basic and the 12-silico-acids are four-basic. Also that the 12-acids and their salts show a marked tendency to crystallize with cubic symmetry, which indicates a highly symmetrical molecule. The proposed molecule is a co-ordinated structure in which 12 tungsten atoms, each surrounded by an octahedron of 6 oxygen atoms, are linked in a continuous shell, by sharing oxygen atoms, round a central  $PO_4$  or  $SiO_4$  tetrahedral group. Each  $WO_6$  octahedron shares three oxygen atoms with neighbouring  $WO_6$  octahedra. In the complete acidic anion there are 18 oxygen atoms, each

\* 'Pogg. Ann.,' vol. 6, p. 369 (1826)

† 'C. R. Acad. Sci. Paris,' vol. 55, p. 838 (1862)

‡ 'Z. Naturwiss. Halle,' vol. 40, p. 298 (1872)

§ 'Bull. Soc. Chim.' vol. 36, p. 221 (1881)

|| 'Z. anorg. Chem.,' vol. 12, p. 16 (1896)

¶ 'Ber. deuts. chem. Ges.,' vol. 20, p. 1811 (1887)

\*\* 'C. R. Acad. Sci. Paris,' vol. 92, p. 1234 (1881)

†† 'Ann. Chim. Phys.,' vol. 7, p. 118 (1906), 'Bull. Soc. Chim.,' vol. 3, p. 101 (1906).

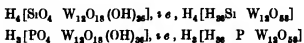
‡‡ 'J. Pract. Chem.,' vol. 77, p. 417 (1906)

§§ Rosenheim and Pinsker, 'Z. anorg. Chem.,' vol. 70, p. 73 (1911), Rosenheim and Jaenicke, 'Z. anorg. Chem.,' vol. 100, p. 304 (1917), vol. 101, p. 247 (1917).

||| 'The Silicates in Chemistry and Commerce,' London, vol. 15, p. 78 (1913)

¶¶ 'J. Amer. Chem. Soc.,' vol. 51, p. 2368 (1929)

shared between two tungsten atoms, thus having their charges completely satisfied. Each of the remaining 36 oxygen atoms is linked to 1 tungsten atom only, and takes up a hydrogen atom to satisfy its charge, giving 36 OH groups as an outer shell. The symmetry is of the cubic point group  $T_d$ . This gives the formulae



Thus this type of molecule cannot exist for the dehydrated forms of the 12-acids reported, with oxygen contents less than 58.

Scroggie and Clark\* attacked the problem of the structure of 12-silicotungstic acid both by chemical and X-ray methods.

The acid dried at  $100^\circ C$  was examined by X-rays using the powder method. They obtained lines corresponding to a body-centred cube of edge = 12.16 Å. By density measurements they found that the unit cell contained two molecules. No attempt was made to work out the positions of the atoms, or to get further details from this photograph. Their work showed that 12-silicotungstic acid is 4-basic. The acid dried at  $100^\circ C$  they represent by  $4H_2O \cdot SiO_2 \cdot 12WO_3 \cdot 4H_2O, \cdot e, H_{16}SiW_{12}O_{44}$ .

This form of the acid they found to be very persistent, and very strong dehydrating conditions were necessary to remove more water. They succeeded in dehydrating to  $2H_2O \cdot SiO_2 \cdot 12WO_3, \cdot e, H_4SiW_{12}O_{40}$ , and found that this apparently was the anhydrous acid, as the removal of any further oxygen caused the acid to change its physical and chemical properties completely, indicating a breakdown of the molecule. From these results they proposed a molecule for 12-silicotungstic acid which gave the formula of 12-silicotungstic acid as  $H_4[H_{12}SiW_{12}O_{44}]$ .

M and E Kahane† studied the dehydration of 12-phosphotungstic acid and came to the conclusion that the dehydrated acid has a formula of the type  $H_nP W_{12}O_{40}$ . As the classical Rosenheim-Miolati formula and also Pauling's proposed formula have more than 40 oxygen atoms in the molecule, they consider these formulae incompatible with their results for dehydrated forms of the acid, and further, consider it unlikely that there should be a different molecule for more highly hydrated forms of the acid.

A large number of hydrates of the 12-acids and their salts have been described by many workers. Representing the 12-acids by the typical formula

\* 'Proc. Nat. Acad. Sci. Wash.' vol. 16, p. 1 (1929)

† 'Bull. Soc. Chim.', vol. 49, p. 557 (1931).

$H_nRM_{12}O_n$ , including water of crystallization to avoid confusion, hydrates have been described where  $n = 40, 44, 46, 58, 58\frac{1}{2}, 61, 62, 65, 68, 69, 69\frac{1}{2}, 70, 70\frac{1}{2}$

In addition to the normal 12-acids which have been discussed, so-called iso-12-acids have been reported in the cases of 12-silicotungstic and 12-borotungstic acid. The cation-cation ratio is reported to be still 1-12, but the properties of these iso-12-acids differ from those of the normal 12-acids, the iso-acids and their salts crystallize in more complex forms with different amounts of water of crystallization, etc. Iso-12-silicotungstic acid was first reported by De Marignac,\* who also prepared its salts. Copaux† and Rosenheim‡ have also studied these iso-12-acids. The iso-12-acids are thought to be isomeric forms of the normal 12-acids. The literature, however, is not very definite concerning these iso-acids. Rosenheim, in his proposed formula, attempts to account for their existence by assigning different properties to different atomic positions in his molecule, whilst Pauling suggests that they may be explained by assuming condensation of two molecules of the normal 12-acid to form the molecule of the iso-12-acid, which would then contain 24 tungsten atoms and 2 silicon or boron atoms.

### 3 Discussion of Literature

The literature concerning the 12-heteropolyacids is very large and much confusion exists owing to uncertainty as to the correct formula of the molecule. Many formulæ have been suggested which differ widely. The classical chemical formula attributes a basicity of 7 to the 12-phospho-acids and a basicity of 8 to the 12-silico-acids, but the experimental evidence for this is not conclusive. Only in very rare cases have salts been reported in which this number of hydrogen atoms have been replaced, *e.g.*, some Ag and guanidine salts, and even for these salts it is more normal to have only 3 and 4 hydrogens replaced instead of 7 and 8 respectively. The salts in which 7 and 8 hydrogens are replaced are only prepared with difficulty. The electrometric evidence supporting these high basicities again cannot be considered conclusive. Practically all salts reported support the formulæ which treat with 12-phospho-acids as 3-basic and the 12-silico-acids as 4-basic, and modern workers tend to consider these basicities as correct. Also the reactions of the 12-acids indicate a structure of  $MO_4$  octahedra rather than one of  $MO_4$  tetrahedra. For example,

\* 'Ann. Chim. Phys.' vol. 3, p. 5 (1864)

† 'Ann. Chim. Phys.' vol. 17, p. 217 (1909)

‡ Rosenheim and Jasenick, 'Z. anorg. Chem.' vol. 101, p. 235 (1917)

the  $\text{MoO}_4$  group is not usually highly coloured, whereas 12-phosphomolybdic acid is of a bright orange colour, which colour is a characteristic of the  $\text{MoO}_4$  group. Again, the fact that these acids are very susceptible to reduction would seem to suggest  $\text{MO}_4$  groups rather than  $\text{MO}_3$  groups.

References to hydrates are meaningless unless the formula used for the acid is given. Some workers tend to treat different hydrates of the same type of acid as distinct acids, describe them as coming out of the same solution together, with differing crystalline forms, and from these differing crystals prepare salts which they declare to differ according to the hydrate of the acid used. Some of these statements are due possibly to the use of impure heteropolyacids, great care must be taken in preparing the 12-acids to ensure that no 9-acid is present. Throughout all the published work, however, there appears to be general agreement that the hydrate of the 12-acids which crystallizes out of saturated aqueous solution at room temperature is of the type  $\text{H}_2\text{RM}_{12}\text{O}_{70}$ , though some workers suggest  $\text{H}_2\text{RM}_{12}\text{O}_{69}$ , and also that when salts of these acids are dehydrated, the lowest form of the anion is  $\text{RM}_{12}\text{O}_{40}$ , removal of more oxygen results in a complete change of properties. There is also much evidence that when 12-acid is dehydrated the lowest form is  $\text{H}_2\text{RM}_{12}\text{O}_{40}$ . In fact several formulæ have been proposed which are based on these facts, the molecule being represented as  $\text{H}_2\text{RM}_{12}\text{O}_{40}$ .

#### 4 The Results of the Present Work

*A General Account of the Structure of the Molecule of 12-phosphotungstic Acid, with its Formula, as found by X-ray Analysis*\*—The derivation of this structure is given in a later section of this paper. The molecule of 12-phosphotungstic acid has the formula  $\text{H}_2[\text{PW}_{12}\text{O}_{40}]$ . The phosphorus atom is at the centre of a group of 4 oxygen atoms arranged with their centres at the corners of a regular tetrahedron as in fig. 1. Each tungsten atom is approximately at the centre of a group of 6 oxygen atoms, the centres of which are at the corners of a distorted octahedron, as in fig. 2.

The complete acidic anion  $\text{PW}_{12}\text{O}_{40}^{-3}$  is a co-ordinated structure of point group symmetry  $T_d$ , consisting of a central  $\text{PO}_4$  tetrahedron, surrounded by 12  $\text{WO}_4$  octahedra as a shell, linked together by shared oxygen atoms. The 12  $\text{WO}_4$  octahedra are arranged in four groups. Each group consists of 3  $\text{WO}_4$  octahedra round a trigonal axis, so that the co-ordinates of the 3 tungsten atoms are of the type  $(a \ a \ b)$   $(a \ b \ a)$   $(b \ a \ a)$ , with two co-ordinates

\* See also 'Nature,' vol. 131, p. 908 (June 24, 1933).

equal, referred to the rectangular axes of cubic symmetry, with origin at the centre of the central phosphorus atom. Fig 3 shows the arrangement for one group of 3  $\text{WO}_6$  octahedra relative to the  $\text{PO}_4$  tetrahedron. Considering first only this one group of three octahedra, there is 1 oxygen atom shared in

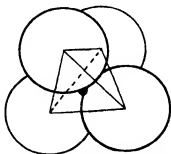


FIG 1—Four oxygen atoms arranged with centres at the corner of a tetrahedron with a phosphorus atom at the centre

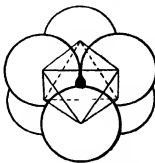


FIG 2—Six oxygen atoms arranged with centres at the corners of an octahedron, with a tungsten atom at the centre

common between the 3  $\text{WO}_6$  octahedra and the central  $\text{PO}_4$  tetrahedron and each octahedron shares two other oxygen atoms, one with each of the two neighbouring octahedra. Thus each octahedron has two edges shared, one

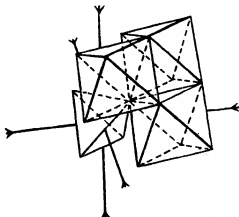


FIG 3—The arrangement of one group of three  $\text{WO}_6$  octahedra relative to the central  $\text{PO}_4$  tetrahedron.

with each of the two neighbouring octahedra. Fig 4 shows this arrangement exploded outwards for simplicity. The arrows indicate the corners which coincide when in their correct positions as shown in fig 3

Four such groups of 3  $\text{WO}_6$  octahedra are arranged round the central  $\text{PO}_4$  tetrahedron in tetrahedral symmetry, so that each oxygen atom of the  $\text{PO}_4$  tetrahedron is now an oxygen atom already shared between 3  $\text{WO}_6$  octahedra. Fig. 5 shows the arrangement with two groups of 3  $\text{WO}_6$  octahedra. For a

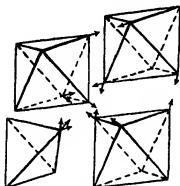


FIG. 4—Exploded diagram of the arrangement shown in fig. 3

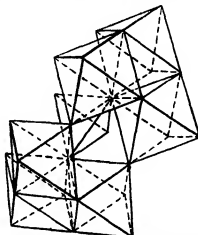


FIG. 5—The arrangement of two groups of 3  $\text{WO}_6$  octahedra relative to the central  $\text{PO}_4$  tetrahedron.

diagram of the complete acidic anion, in order to avoid confusion, each group of 3  $\text{WO}_6$  octahedra is represented in outline as a solid as in fig. 6, which is a representation of the 3 octahedra in fig. 3. The complete acidic anion is shown in fig. 7, the central  $\text{PO}_4$  tetrahedron being shown complete. It will be seen that each  $\text{WO}_6$  octahedron, besides sharing 2 oxygen atoms with octahedra in its own group of three, also shares 2 oxygen atoms with octahedra in other groups.

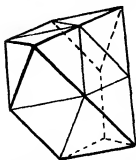


FIG. 6—Simplified representation of the 3  $\text{WO}_6$  octahedra shown in fig. 3

In the complete acidic anion, each  $\text{WO}_6$  octahedron consists of 1 oxygen atom shared between 3  $\text{WO}_6$  octahedra and 1  $\text{PO}_4$  tetrahedron, 4 oxygen atoms shared between 2  $\text{WO}_6$  octahedra, and 1 oxygen atom unshared with other polyhedra. Also, two edges of each  $\text{WO}_6$  octahedron are shared with edges of two other octahedra.

This anion has the formula  $\text{PW}_{12}\text{O}_{40}$ .<sup>3</sup> Hence the molecule of 12-phosphotungstic acid has the formula  $\text{H}_3\text{PW}_{12}\text{O}_{40}$ . This might be written  $\text{H}_3[\text{PO}_4(\text{WO}_6)_3]_4$ , to indicate a central  $\text{PO}_4$  tetrahedral group surrounded by 12  $\text{WO}_6$  octahedra, each of which is  $\text{WO}_3$  in effect. But perhaps

the best form of expression is  $H_3[P(W_3O_{10})_4]$ , to indicate a modified  $PO_4^{-3}$  tetrahedral group, in which each of the 4 oxygen atoms has been replaced by

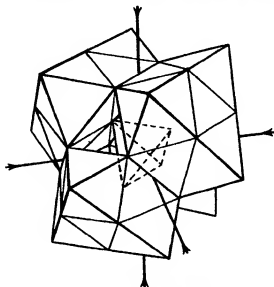


FIG. 7.—The complete anion  $PW_{12}O_{40}$ .

a  $W_3O_{10}^{-2}$  group, each  $W_3O_{10}^{-2}$  group consisting of 3  $WO_6$  octahedra sharing oxygen atoms with other octahedra. This formula agrees with the mass of chemical evidence.

*Co-ordinates of Atoms, and Interatomic Distances*

—The oxygen atoms fall into four groups, which will be called  $O_1$ ,  $O_2$ ,  $O_3$  and  $O_4$ . The positions of these types of oxygen atoms are indicated in fig. 8, which represents the left-hand octahedron in fig. 3. The  $O_1$  oxygen atoms are shared between 3  $WO_6$  octahedra and 1  $PO_4$  tetrahedron. The  $O_2$  oxygen atoms are shared between 2  $WO_6$  octahedra. The  $O_3$  oxygen atoms are also shared between 2  $WO_6$  octahedra. The  $O_4$  oxygen atoms are unshared with other polyhedra. The  $O_1-O_2$  edges are shared between 2  $WO_6$  octahedra.

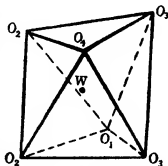


FIG. 8.—The left-hand octahedron in fig. 3, showing the position of the four types of oxygen atoms.

The atoms are referred to the rectangular axes of cubic symmetry, with origin at the centre of the central phosphorus atom.



The co-ordinates of the atoms are —

1 P atom at (0, 0, 0)

4 O<sub>1</sub> atoms at ( $a$ ,  $a$ ,  $a$ ) ( $a$ ,  $\bar{a}$ ,  $\bar{a}$ ) ( $\bar{a}$ ,  $a$ ,  $\bar{a}$ ) ( $\bar{a}$ ,  $\bar{a}$ ,  $a$ )

where  $a = 0.99$  Å

12 O<sub>2</sub> atoms at ( $\bar{b}$ ,  $b$ ,  $c$ ) ( $\bar{b}$ ,  $\bar{b}$ ,  $\bar{c}$ ) ( $b$ ,  $b$ ,  $\bar{c}$ ) ( $b$ ,  $\bar{b}$ ,  $c$ )

( $\bar{b}$ ,  $c$ ,  $b$ ) ( $\bar{b}$ ,  $\bar{c}$ ,  $\bar{b}$ ) ( $b$ ,  $c$ ,  $\bar{b}$ ) ( $b$ ,  $\bar{c}$ ,  $b$ )

( $\bar{c}$ ,  $b$ ,  $b$ ) ( $\bar{c}$ ,  $\bar{b}$ ,  $\bar{b}$ ) ( $c$ ,  $b$ ,  $\bar{b}$ ) ( $c$ ,  $\bar{b}$ ,  $b$ )

where  $b = 0.97$  Å,  $c = 2.84$  Å

12 O<sub>3</sub> atoms at ( $d$ ,  $d$ ,  $e$ ) ( $d$ ,  $\bar{d}$ ,  $\bar{e}$ ) ( $\bar{d}$ ,  $d$ ,  $\bar{e}$ ) ( $\bar{d}$ ,  $\bar{d}$ ,  $e$ )

( $d$ ,  $e$ ,  $d$ ) ( $d$ ,  $\bar{e}$ ,  $\bar{d}$ ) ( $\bar{d}$ ,  $e$ ,  $\bar{d}$ ) ( $\bar{d}$ ,  $\bar{e}$ ,  $d$ )

( $e$ ,  $d$ ,  $d$ ) ( $e$ ,  $\bar{d}$ ,  $\bar{d}$ ) ( $\bar{e}$ ,  $d$ ,  $\bar{d}$ ) ( $\bar{e}$ ,  $\bar{d}$ ,  $d$ )

where  $d = 1.49$  Å,  $e = 3.54$  Å

12 O<sub>4</sub> atoms at ( $\bar{f}$ ,  $g$ ,  $g$ ) ( $\bar{f}$ ,  $\bar{g}$ ,  $\bar{g}$ ) ( $f$ ,  $g$ ,  $\bar{g}$ ) ( $f$ ,  $\bar{g}$ ,  $g$ )

( $\bar{g}$ ,  $f$ ,  $g$ ) ( $\bar{g}$ ,  $\bar{f}$ ,  $\bar{g}$ ) ( $g$ ,  $f$ ,  $\bar{g}$ ) ( $g$ ,  $\bar{f}$ ,  $g$ )

( $\bar{g}$ ,  $g$ ,  $f$ ) ( $\bar{g}$ ,  $\bar{g}$ ,  $\bar{f}$ ) ( $g$ ,  $g$ ,  $\bar{f}$ ) ( $g$ ,  $\bar{g}$ ,  $f$ )

where  $f = 0.1$  Å,  $g = 3.79$  Å

12 W atoms at ( $h$ ,  $k$ ,  $l$ ) ( $h$ ,  $\bar{l}$ ,  $\bar{l}$ ) ( $\bar{h}$ ,  $k$ ,  $\bar{k}$ ) ( $h$ ,  $\bar{k}$ ,  $l$ )

( $k$ ,  $h$ ,  $\bar{k}$ ) ( $k$ ,  $\bar{h}$ ,  $\bar{l}$ ) ( $\bar{l}$ ,  $h$ ,  $\bar{k}$ ) ( $\bar{k}$ ,  $\bar{h}$ ,  $k$ )

( $k$ ,  $k$ ,  $h$ ) ( $k$ ,  $\bar{l}$ ,  $\bar{h}$ ) ( $\bar{k}$ ,  $k$ ,  $\bar{h}$ ) ( $\bar{k}$ ,  $\bar{k}$ ,  $h$ )

where  $h = 0.07$  Å,  $k = 2.495$  Å

The interatomic distances are as follows —

The PO<sub>4</sub> tetrahedron in the centre is undistorted. All edges (O<sub>1</sub>-O<sub>1</sub> distances) are 2.80 Å.

The WO<sub>4</sub> octahedra are distorted. The shared edges (O<sub>1</sub>-O<sub>2</sub> distances) are 2.65 Å. The O<sub>1</sub>-O<sub>2</sub> distances are 2.70 Å. The O<sub>2</sub>-O<sub>2</sub> distances are 2.90 Å.

The O<sub>2</sub>-O<sub>2</sub> distances are 2.65 Å. The O<sub>2</sub>-O<sub>2</sub> distances are 2.61 Å. The O<sub>2</sub>-O<sub>4</sub> distances are 3.10 Å. The O<sub>4</sub>-O<sub>2</sub> distances are 2.80 Å.

The tungsten atom is 2.34 Å distant from the O<sub>1</sub> oxygen atom, 1.85 Å distant from the O<sub>2</sub> and O<sub>4</sub> oxygen atoms and 2.0 Å distant from O<sub>3</sub> oxygen atoms.

## 5 Discussion of this Structure

From a consideration of the forces acting in this structure, these distances and the distortions are seen to be quite logical. The average crystal radii of the P, W, and O atoms may be taken as equal to 0.34 Å, 0.62 Å, and 1.35 Å respectively. The tungsten atoms have a major effect in determining the structure, and the positions of the oxygen atoms will be to a large extent governed by them. The tungsten atoms repel each other, and tend to distribute themselves in the most uniform manner possible where their potential energy is lowest. Neglecting the very small displacement (0.07 Å) of these atoms from the ideal positions, the reasons for which will be dealt with later, in this structure this distribution is realized, the 12 tungsten atoms are arranged on a sphere with the most uniform distribution possible, each tungsten atom being equidistant (3.53 Å) from four neighbouring tungsten atoms. They may also be regarded as being at the centres of the edges of a regular octahedron, or at the centres of the edges of a cube. The oxygen atoms then form distorted octahedra round these tungsten atoms in such a manner as to give this distribution for minimum potential energy of the tungsten atoms most closely. The edge of a  $\text{PO}_4$  tetrahedron is normally about 2.6 Å. In this case the edge is lengthened to 2.8 Å, as 3 tungsten atoms act in opposition to the central phosphorus atom and tend to pull the oxygen atoms of the  $\text{PO}_4$  group outwards. The tungsten atoms are not at the centres of the octahedra. Mutual repulsion tends to make the sphere on which they lie be of as large a radius as possible, and in this structure this effect is considerable. Hence the distance of the tungsten atoms from the inner  $\text{O}_1$  oxygen atoms is relatively large (2.34 Å) as compared with their distances from the other oxygen atoms. Also, in order to reach positions of minimum potential energy, the tungsten atoms are displaced from the centres of the octahedra so as to be nearer the  $\text{O}_2$  oxygen atoms (1.85 Å distant) than the  $\text{O}_3$  oxygen atoms (2.0 Å distant). In agreement with this displacement, the  $\text{O}_2$ - $\text{O}_3$  distances are lengthened to 2.90 Å and the  $\text{O}_1$ - $\text{O}_2$  distances and the shared edges, ( $\text{O}_1$ - $\text{O}_3$  distances) are shortened to 2.65 Å.

The formula  $\text{H}_2\text{PW}_{12}\text{O}_{40}$  agrees with the large mass of evidence already mentioned, which gives this formula for the anhydrous acid.

This structure has some remarkable features. It appears that some of the principles determining the structure of complex ionic crystals as formulated by Pauling\* are not strictly obeyed. Each of the 4 oxygen atoms of the type  $\text{O}_1$

\* 'J. Amer. Chem. Soc.' vol. 51, p. 1010 (1929)

which form a tetrahedron round the central phosphorus atom, is shared between 1 phosphorus and 3 tungsten atoms. According to Pauling's "electrostatic valence principle," where the electrostatic valence bonds of the cation act in a roughly symmetrical manner and are neutralized by the immediate neighbouring anions, this implies that each  $O_1$  oxygen atom has a force of  $+4\frac{1}{2}$  acting upon it, if the phosphorus atom has charge of  $+5$  and the tungsten atom a charge of  $+6$ . Again, there is an outer shell of 12  $O_4$  oxygen atoms, each of which is directly connected with only 1 tungsten atom. According to the above principle each of these oxygen atoms has a force of only  $+1$  acting upon it. This structure may be considered in two ways. (A) If the bonds of the cations act in a roughly symmetrical manner, then this complex anion has a region of large positive potential at its centre, surrounded by a region of negative potential as a shell, which, by action through a distance across intervening atoms, gives a total charge of  $-3$ , or (B) the electrostatic valence bonds of the cations do not necessarily act in a symmetrical manner, the directions of their action being determined by the positions of the unsatisfied anions.

The properties to be expected from this structure agree well with the known properties of 12-phosphotungstic acid.

This acidic anion would be expected to be stable, as the tungsten atoms are practically in positions of minimum potential energy, and the whole structure is compact and tightly bound together.

These acidic anions are complete structural units in themselves, and when packed together in a crystal are not strongly linked to each other, and should thus be relatively easy to separate. Thus the acid would be expected to be highly soluble.

Also any crystalline structure formed by the packing together of these anions would be expected to contain a relatively large amount of water of crystallization. For these anions are large, roughly spherical units, and, when arranged even in close packing, will still leave a considerable volume of unoccupied space. Water of crystallization will tend to pack in these spaces so as to make the whole structure more homogeneous, and as oxygen atoms are very small compared with these acidic anions, there is always room for a relatively large amount of water.

This anion also explains why numerous hydrates are formed by the acid. As the atoms are arranged in a fairly uniform manner, and the charge is small for such a large group of atoms, this anion may be considered as a large, practically spherical, unit, with a fairly uniform charge distribution over its

surface. Such a unit has no strong directional forces which would tend to make one type of packing more probable than any other arrangement. Thus it is to be expected that quite a number of arrangements would exist. Also, as these units are so uniform in shape and charge distribution, they will tend to pack in simple arrangements which explains the tendency for the hydrates to be cubic, despite the complexity of the anion.

The tungsten atom, by reason of its small size, may be surrounded by a tetrahedron of oxygen atoms, as well as by an octahedron, as in this structure. Thus the reduction of the present arrangement should not be difficult.

The marked tendency of these acids to combine with large organic molecules to form insoluble precipitates is possibly connected with the large, shell-like region of negative potential surrounding the anion.

The basicity of three for this structure agrees with that found by most chemical workers. Also the oxygen content of the molecule—40 atoms of oxygen—agrees exactly with the results obtained for the anhydrous acid by many workers on the dehydration of the acid and its salts.

## 6 Experimental

The preparation of the acid used in this work was that described by Wu.\* It was found that if no HCl was added during the last ether extraction, the product was pure 12-phosphotungstic acid, though the omission of HCl reduces the yield.

The colourless octahedra which crystallize from aqueous solutions at room temperatures are extremely unstable, and commence to lose water and to break down to a white powder immediately on drying. Thus, though by taking precautions against loss of water an apparently good photograph of this hydrate was obtained, this photograph was not used in this work owing to the uncertainty introduced by the instability of the crystal. The structure of this hydrate, however, has since been determined.

*Dehydration of the Acid*—In order to obtain a more definite photograph, a more stable hydrate was sought. It was assumed that the hydrate which crystallizes out of saturated aqueous solution at room temperature in colourless octahedra was of empirical formula either  $H_2PW_{12}O_{70}$  or  $H_2PW_{12}O_{66}$ . This was the only assumption made as regards the formula, and losses of water on dehydration were calculated on the first formula. Whether  $H_7$  or  $H_3$  is

\* 'J. Biol. Chem.', vol. 43, p. 199 (1920).

taken as the basicity is immaterial, owing to the large molecular weight of the acidic anion

In order to find the loss in weight, it was necessary to weigh this unstable hydrate To do this with as little error as possible, large crystals were grown, of dimensions of the order of 1 cm These crystals were taken out of the solution, dried quickly with filter paper, put in a closely stoppered bottle and weighed immediately By using large crystals, surface action had a relatively small effect and it was found possible to repeat results quite accurately

This unstable hydrate was allowed to reach constant weight in the open at about 70% RH 15.6 mols of water were lost An X-ray photograph of this product showed a mixture with no distinct lines

Drying over  $\text{CaCl}_2$  at atmospheric pressure and room temperature was next tried The acid reached constant weight with a loss of 22.5 mols of water On standing in the open this product took up only 0.5 mols of water, which was probably due to surface adsorption, and indicated that this product is fairly stable This gave a photograph with sharp lines On drying over  $\text{P}_2\text{O}_5$  in *vacuo*, the weight was practically constant with a loss of 23.5  $\text{H}_2\text{O}$  On prolonged drying, however, over a period of several days, the weight continued to fall very slowly, until after about a week, 24 mols of water were lost After this period the acid commenced to turn brown on the surface, indicating that disintegration of the molecule had commenced For losses of water ranging from 22.5 mols to 23.5 mols the product gave identical sharp lines but after losses of more than 23.5 mols of water the lines commenced to broaden, and after a loss of 24 mols of water the lines were distinctly blurred, which indicates the beginning of a change in the crystal structure There is a definite structure in this range, but the number of molecules of water present is not definitely fixed by the above results The water content of this hydrate will be considered later

*X-ray Technique*—The X-ray photographs were taken by the powder method in a camera of the modified Debye-Scherrer type as described by Bradley and Jay\* The specimen to be photographed is in the form of a powder rod Owing to the chemical activity of this acid, and its variable water content, the normal method of preparing the powder rod, by fixing the powder to a hair with adhesive, was not used In order to avoid the possibility of change, the powder was packed into a fine capillary of Landemann glass, which was sealed, mounted in the camera and used as a specimen. This glass

\* 'Proc Phys. Soc.,' vol 44, p. 563 (1932)

has a low absorption for X-rays, being composed of lithium and beryllium borates

The observed intensities of reflection were obtained by measuring the blackening of the powder photograph by means of a Cambridge microphotometer designed by Dobson\*. The figures for blackening were then converted into intensity values by means of a calibration curve for the emulsion used, in which exposure is plotted against blackening. The intensity measurements were then plotted against distance along the film, and an integrated value of the intensity of each line was obtained by determining the area under the curve and above the base line of general blackening.

The structure investigated in this work was that of the dehydrated 12-phosphotungstic acid discussed before, which gave sharp lines. The acid actually photographed was the product obtained by dehydrating the octahedral crystals over  $P_2O_5$  *in vacuo*, with a loss in weight corresponding to loss of 23.5 mols. of water.

$CuK_\alpha$  radiation was used. A typical powder photograph of this hydrate is shown in Plate I.

### 7 Determination of the Lattice

The values of  $\Sigma h^2$  for the lines were found to be all even numbers, except for two very faint lines corresponding apparently to  $\Sigma h^2 = 33$  and 41. These two lines were neglected at first, as it was thought that they were due to slight impurity. An arrangement of lines where  $\Sigma h^2$  corresponds to consecutive even numbers is a characteristic of the body-centred cubic arrangement. Also, as lines where  $\Sigma h^2 = 28, 60, 92$ , etc., which are theoretically impossible, were missing from the photograph, it was assumed, at first, that this structure was a body-centred cubic arrangement. It was found later that this was not quite the true arrangement. The effect of the modification necessary, is, however, extremely small. This will be considered later. The value of " $a$ ," the length of the edge of the unit cube, was now calculated for each line. The results are given in Table I.

The usual practice of comparing the observed  $\sin^2 \theta$  with the calculated  $\sin^2 \theta$  is unnecessary here.

By plotting these values of " $a$ " against  $\cos^2 \theta$ , and extrapolating to  $\cos^2 \theta = 0$ ,† the true value of the edge of the unit cube was found to be  $12.141 \pm 0.005 \text{ \AA}$ .

\* 'Proc. Roy. Soc.,' A, vol. 104, p. 246 (1923).

† 'Proc. Phys. Soc.,' vol. 44, p. 563 (1932).

The density of this hydrate was determined by the displacement of benzene, using a density bottle, and was found to be 5.6. The density calculated for two molecules of  $H_2PW_{12}O_{40}$  in a cube of side 12.14 Å is 5.55. This agrees well with the experimental value. For a body-centred cubic arrangement, this implies that there is an anion at each corner of the unit cube and one at the centre, all anions being equivalent for this symmetry.

Table I

$\Sigma h^2$	"a" in Å	$\Sigma h^2$	"a" in Å	$\Sigma h^2$	"a" in Å
2	11.665	60	—	118	—
4	11.827	62	12.113	120	—
6	11.891	64	—	122	12.139
8	11.937	66	12.116	124	—
10	11.955	68	12.118	126	12.141
12	11.981	70	12.122	128	—
14	12.004	72	12.122	130	—
16	12.007	74	12.121	132	12.147
18	12.027	76	—	134	12.145
20	12.035	78	12.124	136	—
22	12.040	80	—	138	12.147
24	—	82	12.125	140	—
26	12.052	84	—	142	12.144
28	—	86	12.129	144	—
30	12.069	88	—	146	—
32	12.080	90	12.130	148	—
34	12.079	92	—	150	12.147
36	12.078	94	12.127	152	12.143
38	12.075	96	—		
40	12.083	98	12.130		
42	12.083	100	12.129		
44	12.085	102	12.129		
46	—	104	—		
48	12.093	106	12.134		
50	12.101	108	—		
52	12.106	110	12.132		
54	12.099	112	—		
56	12.101	114	—		
58	12.111	116	12.135		

The above, together with the observed intensity values, were all the data available, as no single crystal data could be obtained.

The problem was now to locate the atoms in the structure by use of the above data.

### 8 Location of the Tungsten Atoms

As the scattering power of a tungsten atom for X-rays is much greater than that of either an oxygen or a phosphorus atom, it was possible to locate the 12 tungsten atoms quite accurately, by neglecting all other types of atoms at first, and considering the agreement between the observed intensities and the

intensities calculated for the 12 tungsten atoms only. It was assumed that there were 12 tungsten atoms in each molecule of the acid, and that the anions were arranged in body-centred cubic packing.

The simplest way to arrange groups of 12 tungsten atoms to agree with body-centred cubic symmetry, is to arrange them in the manner already given in the section describing the structure of the anion, each group of 12 tungsten atoms having point group symmetry  $T_d$ . In this arrangement the positions of the atoms are defined by only two variables, and it is thus possible to represent atomic positions on a graph. When such groups of atoms are arranged with body-centred packing, the space group is  $T_d^3$ . From a consideration of the observed intensities it was possible to impose very rough limits for the calculated intensities which would be consistent with those observed on the film. In this way curves defining areas of possible atomic positions could be drawn on a graph for each reflection. This was done for a large number of reflections until only a small area of possible positions for the tungsten atoms was left.

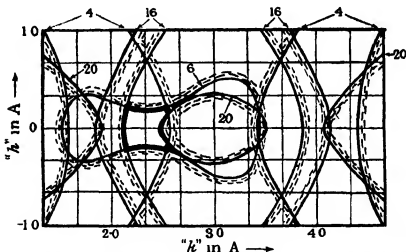


FIG. 9—Curves locating the final small area of possible positions for the tungsten atoms. The final area is heavily outlined.

Fig. 9 shows the location of the final area of possibility. This area is shown heavily outlined, and the reflections used for this location are shown. The areas on the shaded sides of the curves are areas of forbidden co-ordinates for the tungsten atoms. The area left gives possible co-ordinates for the tungsten atoms between the limits  $h = 2.155 \text{ Å} \rightarrow 2.51 \text{ Å}$ ,  $h = 0.185 \text{ Å} \rightarrow 0 \rightarrow -0.185 \text{ Å}$ . Thus the possible variation of  $h$  or  $k$  is not more than about  $0.36 \text{ Å}$ . The location is



surprisingly definite, even though the limits of possible intensities imposed were quite rough, owing to the large allowances made for the possible effect of atoms other than tungsten. In spite of this, a sufficient number of sensitive lines, for which the intensity varies very rapidly for small changes in the atomic co-ordinates, were found to give this small final area of possibility. A similar method was used by W. L. Bragg and Warren\* in work on the silicates. The mean location is  $h = 0$ ,  $k = 2.33$  Å.

To get a still more accurate location of the tungsten atoms, the intensities of many lines were now worked out for sets of co-ordinates in this area, still considering only the tungsten atoms. By considering the best general agreement between observed and calculated intensities, it appeared that the most probable co-ordinates for the tungsten atoms are  $h = 0$ ,  $k = 2.495$  Å. The general agreement for these co-ordinates is reasonably good for all lines.

#### *9 The Structure of Co-ordinated Oxygen Polyhedra agreeing with these Tungsten Co-ordinates*

It was assumed that the oxygen atoms would be arranged round the phosphorus and tungsten atoms with their centres at the corners of polyhedra, the cations being roughly at the centres of these polyhedra, as is usual for structures of the complex ionic type.

The crystal radii of the phosphorus, tungsten and oxygen atoms are usually taken as equal to 0.34 Å, 0.62 Å, and 1.35 Å respectively.

The positions of the tungsten atoms were fixed, and to agree with symmetry a phosphorus atom must be at the centre of each group of 12 tungsten atoms. An arrangement of oxygen polyhedra around these fixed atoms had now to be found which would fulfil the following conditions —

- (1) The distances between the centres of neighbouring oxygen atoms in the structure must be within the limits 2.50 Å to 3.10 Å.
- (2) The acidic anions must pack in a cube of side = 12.14 Å so that the distances between the centres of neighbouring atoms in adjacent anions are of the correct dimensions. There must be two anions in each cube.
- (3) The number of hydrogen atoms necessary to give a neutral structure, i.e., the basicity of the acid, must be reasonable, preferably 3 or 7 to agree with the bulk of chemical evidence.
- (4) The stability of the acid suggests that the polyhedra should be well linked together by sharing oxygen atoms.

\* 'Z. Kristallog.' vol. 69, p. 168 (1923).

Many arrangements were tried with various types of polyhedra, but the conditions imposed were found to be very stringent, and condition 1 alone eliminated decisively all arrangements other than the anion which was found to be correct and which has been described. This structure was found to fulfil all the other conditions, and gave the formula of the molecule as  $H_2PW_{12}O_{40}$ . It is interesting to note that in none of the structures tried was a basicity of 7 indicated for this acid.

# 10 The Arrangement of these Anions in the Structure of the Hydrate Investigated.

The symmetry of this anion is essentially tetrahedral, and in considering the orientation of the anions it is convenient to refer to the orientation of the central  $PO_4$  tetrahedron. The  $O_3$  oxygen atoms are above a base, and the  $O_4$  oxygen atoms are above an apex of this central tetrahedron.

The structure which has been described for the anion of 12-phosphotungstic acid is very symmetrical. At first it was thought that the anions were packed

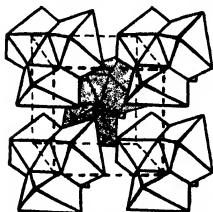


FIG 10—True body centred arrangement ( $T_h^2$ )

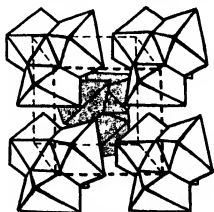


FIG 11—Arrangement  $O_h^4$

in a body-centred cubic arrangement, as all the lines on the powder photograph indicated this type of packing, with the exception of two very faint lines corresponding to  $\Sigma h^2 = 33$  and 41, which were at first neglected as being due to impurity. In this arrangement the orientation of all anions is identical, the anions being arranged base to apex right through the structure. This arrangement is shown in fig 10. The space group for this arrangement is  $T_h^2$ , and no reflections for which  $\Sigma h^2$  is odd are possible.

On further consideration, it was realized, however, that an arrangement where the anions were base to base and nose to nose in the structure would differ only slightly from the true body-centred arrangement. This arrangement is shown in fig. 11. The space group here is  $O_h^4$ , and the only difference between this packing and the previous body-centred packing is that the anion in the centre of the cube is inverted. Thus, if the tungsten co-ordinates are actually  $h = 0$ ,  $k = 2.495 \text{ \AA}$ , there will be no change at all in the arrangement of the tungsten atoms, and as the oxygen arrangement is also fairly symmetrical, the change in the oxygen lattice will not be great. Thus it is to be expected that the photographs for these two types of arrangement would be very similar.

For the  $O_h^4$  arrangement, lines where  $\Sigma h^2$  is odd are possible, but only oxygen atoms contribute to these reflections if  $h = 0$  for the tungsten atoms. Hence these lines will be very weak.

The intensities of reflection for both arrangements were calculated, and compared with the observed intensities, in order to decide which arrangement is correct. An attempt was also made to find positions for the molecules of water of crystallization.

*The True Body-centred Arrangement ( $T_d^2$ )*—In this arrangement the distance between the centres of neighbouring oxygen atoms of adjacent anions is  $2.7 \text{ \AA}$ , which is reasonable. Here the  $O_3$  oxygen atoms of one anion touch the  $O_4$  oxygen atoms of the adjacent anion.

It is impossible to pack six molecules of water\* in this structure to agree with symmetry.

A possible arrangement giving seven molecules of water per molecule of acid may, however, be suggested. A molecule of water may be placed at the centre of each face of the unit cube—at  $b\bar{b}00$ ,  $0b\bar{b}0$ ,  $00b\bar{b}$ ,  $\bar{b}00b$ ,  $0\bar{b}0b$ ,  $00\bar{b}b$ , where  $b = 6.07 \text{ \AA}$ . This gives three molecules of water per molecule of acid. The centres of water molecules in such positions are  $3.2 \text{ \AA}$  from the centres of neighbouring oxygen atoms in the cube face, and  $3.3 \text{ \AA}$  from the centres of neighbouring oxygen atoms in a direction perpendicular to the cube face. These water molecules have thus considerable freedom of motion. The other four molecules of water might be arranged between the acidic anions, on the trigonal axes of the cube, in tetrahedral symmetry, in positions  $aaa$ ,  $a\bar{a}\bar{a}$ ,  $\bar{a}a\bar{a}$ ,  $\bar{a}\bar{a}a$ , where  $a = 3.28 \text{ \AA}$ . Here, however, the centre of each water molecule is only

\* The term "water molecule" is used here in a general sense to indicate those oxygen atoms which could be removed as water without affecting the chemical properties of the acid. No attempt is made to define the positions of the hydrogen atoms.

2.57 Å distant from the centres of six neighbouring oxygen atoms. The distance is somewhat small.

*The Arrangement  $O_h^4$* —In this arrangement, where the two adjacent anions are base to base the oxygen atoms of the anion do not touch, being about 3.5 Å apart. There is, however, just room for a molecule of water in the hollow left here between the anions, which would hold them apart. The water— $O_2$  oxygen distance is 2.9 Å. The positions of these four water molecules are  $\bar{a}\bar{a}\bar{a}$ ,  $\bar{a}aa$ ,  $a\bar{a}\bar{a}$ ,  $aaa$ , where  $a = 3.03$  Å. This arrangement contributes two molecules of water per molecule of acid. Where the two adjacent anions are apex to apex, the  $O_2$  oxygen atoms of one anion touch the  $O_4$  oxygen atoms of the neighbouring anion, the distance between their centres being 2.85 Å. There is no room for a molecule of water between the anions here.

Three more water molecules may be placed, as before, at the centres of the faces of the unit cube. This gives, in all, five molecules of water per molecule of acid for this arrangement.

#### 11 *Comparison of the Intensities of Reflection Calculated for these two arrangements with those observed on the Photograph*

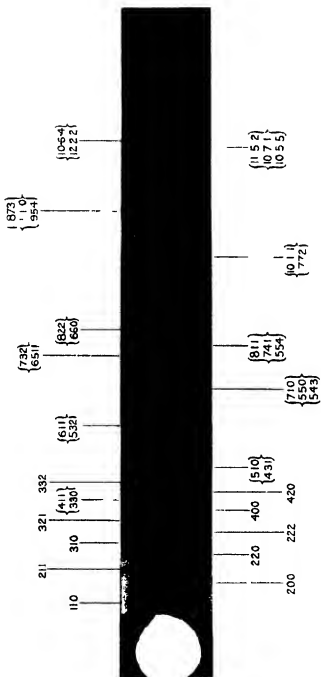
The intensities of reflection for these two arrangements were now calculated and compared with the observed intensities. The results are given in Table II.

For the  $O_h^4$  arrangement, the tungsten atoms were first considered to have co-ordinates  $h = 0$ ,  $k = 2.495$  Å. With such co-ordinates the tungsten atoms do not contribute to reflections where  $\Sigma h^2$  is odd. It was found that, though lines 33 and 41 were the strongest of the odd lines, they were not quite strong enough to account for the observed intensities. It was found, however, that if the tungsten atoms were shifted a very small distance (0.07 Å) to have co-ordinates  $h = 0.07$  Å,  $k = 2.495$  Å, so that these atoms began to contribute to the odd lines, the effect was greatest on lines 33 and 41, and gave calculated values which agreed with the observed intensities.

The third column in Table II gives the calculated intensities of reflection when the tungsten atoms only are considered. The values for lines where  $\Sigma h^2$  is odd refer only to the  $O_h^4$  arrangement, as for the  $T_d^3$  arrangement the values for these reflections are zero. As the calculated intensities for tungsten atoms only, for reflections where  $\Sigma h^2$  is even, are practically identical for the two arrangements, the values given in this column for these reflections refer to both arrangements.

Table II

2A*	AH	Calculated intensities 12 tungsten atoms only for both arrangements	Calculated intensities for T <sub>d</sub> <sup>8</sup> arrangement		Calculated intensities for O <sub>A</sub> <sup>4</sup> arrangement	Observed intensities
			For PW <sub>12</sub> O <sub>40</sub>	For PW <sub>12</sub> O <sub>40</sub> 7H <sub>2</sub> O	For PW <sub>12</sub> O <sub>40</sub> 5H <sub>2</sub> O	
1	100	—	—	—	—	—
2	110	28	46	43	48	43 5
3	111	0 4	—	—	0 6	Absent
4	200	12	13 5	14 5	12 5	12
5	210	—	—	—	—	—
6	211	12	11 5	13	13	12
7	—	—	—	—	—	—
8	220	32 5	42 5	29	32 5	34 5
9	{300 221}	0 3	—	—	0 2	Absent
10	310	28	24	26	26	26 5
11	311	0 2	—	—	0 7	Absent
12	222	87	99	97	101	112
13	320	—	—	—	—	—
14	321	3 5	4	3 2	3	4 1
15	—	—	—	—	—	—
16	400	40	37 5	44 5	42 5	45 5
17	{410 322}	0 01	—	—	0 2	Absent
18	{330 411}	22	18 5	18	18 5	21 5
19	341	0 5	—	—	0 7	Absent
20	420	22 5	17	18	16	13
21	421	0	—	—	0 07	Absent
22	332	116	111	108	109	101
23	—	—	—	—	—	—
24	422	0 02	1	3 1	2	?
25	{500 430}	—	—	—	—	—
26	{510 431}	82	62 5	62 5	62	56 5
27	{511 333}	1 0	—	—	1 6	?
28	—	—	—	—	—	—
29	{520 432}	0 2	—	—	1 1	?
30	521	34 5	17	18 5	18 5	16
31	—	—	—	—	—	—
32	440	15	14	18	16 5	14
33	{441 522}	1 2	—	—	1 8	Just visible
34	{530 433}	6 1	6 3	7 8	6	5 9
35	531	0 5	—	—	0 35	Absent
36	{600 442}	16 5	16 5	17	16	16 5
37	610	—	—	—	—	—
38	{611 532}	39	45	46 5	46	45
39	—	—	—	—	—	—
40	620	14 5	15	10 5	11 5	14



X-Ray powder photograph of 12-phosphotungstic acid over  $P_2O_5$  in *vacuo*  $H_2PW_{12}O_{40}$   $5H_2O$  (n.b., radiation cut  $\lambda = 12.14 \text{ \AA}$ )



Table II—(continued)

Zn*	All	Calculated intensities 12 tungsten atoms only for both arrangements	Calculated intensities for $T_d^*$ arrangement		Calculated intensities for $O_h^*$ arrangement	Observed intensities
			For $PW_{12}O_{40}$	For $PW_{12}O_{40} \cdot 7H_2O$	For $PW_{12}O_{40} \cdot 5H_2O$	
41	{621 540 443}	2 8	—	—	4 15	4
42	541	27	29	27 5	27 5	29
43	533	0 06	—	—	0 35	Absent
44	622	5 6	6 4	6 2	6 8	7 2
45	{630 542}	0 25	—	—	0 5	Absent
46	631	2 1	1 6	1 9	1 9	?
47	—	—	—	—	—	—
48	444	1 9	4	5 2	4 7	5 2
49	{700 632 710 550 643}	0 1	—	—	0 2	Absent
50	{351 711}	98 5	111	112	111	105
51	640	1	—	—	0 8	Absent
52	720	5 6	9 1	8 9	9 6	8 1
53	{641 721 633 552}	0 2	—	—	0 8	Absent
54	—	12 5	14	14	14	11
55	—	—	—	—	—	—
56	642	—	6 7	4	4 3	5 8
57	{722 544}	0 20	—	—	0 0	Absent
58	730	13 5	10	10 5	10 4	10
59	{731 553}	0 45	—	—	0 6	Absent
60	—	—	—	—	—	—
61	{650 643 732 661}	0 1	—	—	0 3	Absent
62	—	117	120	116	118	130
63	—	—	—	—	—	—
64	800	0 16	0 1	0 01	0 2	Absent
65	{810 740 682 811 741 854}	0 35	—	—	1 8	?
66	—	47 5	51	51	51	58
67	733	0 04	—	—	0 2	Absent
68	{820 644}	11	12 2	12 5	12	12 8
69	{821 742}	0 4	—	—	0 95	Absent
70	673	11	11 9	12 2	12 4	12 1
71	—	—	—	—	—	—
72	{822 680}	33	41	46	45	43



All intensities are reduced to the same scale to make comparison simpler. A temperature factor  $e^{-B \sin^2 \theta}$  was applied to the calculated values, where  $B = 2.0$ . In calculating these intensities the usual corrections were made for  $\theta$  and absorption.

No correction was made for the possible effect of motion of the three water molecules in the faces of the cube. Motion would tend to decrease the effect of these molecules on the intensity.

## 12 Discussion of Results

Considering the intensities calculated for the tungsten atoms only, for the lines where  $\Sigma h^2$  is even the agreement with the observed intensities is quite good. This applies to both arrangements. For the odd lines given by the  $O_h^4$  arrangement, the agreement for tungsten atoms only is also reasonable. When the oxygen atoms are included, however, the agreement is much better, for both arrangements. From a consideration of the agreement for the even lines only, it is difficult to decide which arrangement is better, as the differences are very small, though the  $O_h^4$  arrangement would appear to be slightly better.

For the odd lines, however, which are given only by the  $O_h^4$  arrangement, the agreement is also good. In the numerical scale used here to represent intensity values, where strong lines have values in the region of 100, the lower limit of visibility is in the region 1 to 2. Thus line 33 is just visible, and line 41 is distinctly visible. From the calculations it appears that lines 27, 29 and 65 of the odd lines might also be visible, and when looked for could just be detected, though normally they would not be noticed. All the other odd lines were invisible.

This agreement of the odd lines, in addition to a slightly better agreement for the even lines for the arrangement  $O_h^4$ , appears to prove decisively that the anions are arranged in this manner, and not in the body-centred cubic arrangement  $T_d^3$ .

This agreement for the odd lines provides a very sensitive check on the correctness of the proposed structure. For, with 45 oxygen atoms present, it is highly probable that any error in their location would give calculated values for the intensity of some odd reflections which would be well above 2 on this scale, as the effect of the oxygen atoms can be quite large. This is well shown in the various intensity values calculated for lines 2, 26 and 30 in the table.

Thus there are only five molecules of water per molecule of acid in this partially dehydrated acid. This is of particular interest, as later work on the structure of the so-called 30 hydrate of this acid indicates that the octahedral crystals contain only 29 molecules of water per molecule of acid. Most workers have described a loss of 24 molecules of water per molecule of acid when the octahedral crystals are dehydrated to the stable hydrate investigated here. In the present work sharp lines were not obtained for losses of water of more than 23.5 molecules per molecule of acid, but it is highly probable that a certain amount of moisture is adsorbed on the surface of the fine powder. This water, though not actually in the structure, would tend to decrease the loss in weight. Thus a water content of  $5\text{H}_2\text{O}$  for this hydrate may be said to agree with experimental results.

The author wishes to thank Professor W. L. Bragg, F.R.S., for his stimulating interest in the work, which was carried out in the Physical Laboratories of the University of Manchester, also Dr. A. J. Bradley for his helpful discussion.

### 13 Summary

The structure of the molecule of 12-phosphotungstic acid has been found by X-ray analysis, using the powder method. The formula of the acid is  $\text{H}_3[\text{P}(\text{W}_2\text{O}_{10})_4] \cdot 5\text{H}_2\text{O}$ . The anion consists of a central  $\text{PO}_4$  tetrahedral group surrounded by 12  $\text{WO}_6$  octahedral groups as a shell, linked together by shared oxygen atoms. The symmetry is  $T_d$ . The positions of the atoms are given. This structure has some unusual features. In the partially dehydrated acid investigated, which contains five molecules of water per molecule of acid, these acidic anions pack together with cubic symmetry, the edge of the unit cube being  $12.141 \pm 0.005 \text{ \AA}$ . Symmetry  $O_h$ . There are two molecules of acid in the unit cube. This formula agrees with the bulk of chemical evidence.

The structure was worked out rigorously from experimental results. The method used is described. From the observed intensities of reflection the positions of the heavy tungsten atoms were found, in this first stage the effect of all other types of atoms being neglected. A graphical method was used. This preliminary location of the tungsten atoms was surprisingly definite.

A structure of co-ordinated oxygen polyhedra was then found which would give these tungsten positions and also fulfil all other conditions. The com-

parison of observed intensities and those calculated for the complete acid anion is given. The agreement is good and proves the correctness of the structure.

A brief summary of the properties of the 12-heteropoly acids, and the more important papers in the literature relating to their constitution, is given and discussed.

The structure found in the present work is described.

This structure is discussed and its bearing on the properties of the acid is considered.

The preparation of pure 12-phosphotungstic acid is given and its dehydration described.

The experimental X ray technique is described briefly.

The method of locating the tungsten atoms is described and the conditions to be fulfilled by the structure are given. Two types of arrangement of these anions in the unit cube were found to be possible. These are discussed.

The problem of the positions of the molecules of water of crystallization for both types of arrangement is discussed.

The comparison of observed and calculated intensities is given, and discussed.

# *A General Proof of Certain Fundamental Equations in the Theory of Metallic Conduction*

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## § 1 Introduction

In the modern theory† of electronic conduction the electrons are considered, when the thermal motion of the lattice is neglected, as moving in a periodic potential with the property

$$V(x + la, y + ma, z + na) = V(x, y, z)$$

The wave equation for an electron in this field is

$$\left\{ \frac{\hbar^2}{8\pi^2 m} \nabla^2 + E_K - V \right\} \psi_K = 0 \quad (1)$$

Bloch has shown that this equation has solutions of the form

$$\psi_K = e^{i\mathbf{K} \cdot \mathbf{R}} U_K(\mathbf{R}),$$

where  $U_K$  has the periodicity of the lattice

The quantity  $(\hbar/2\pi)\mathbf{K}$  has properties analogous to the momentum of free electrons. In particular, the statistical weight of an element of volume in  $\mathbf{K}$  space is independent of  $\mathbf{K}$ , so that the number of electrons per unit volume of  $\mathbf{K}$  space is constant for all values of  $\mathbf{K}$  having energy below the surface of the Fermi distribution. This surface, which is, of course, identical with a surface of constant energy, is spherical only in the special case of free electrons (fig. 1)

The average velocity of an electron in state  $\psi_K$  is

$$\mathbf{v}_K = \frac{\hbar}{4\pi m i} \{ \psi_K^* \text{grad } \psi_K - \psi_K \text{grad } \psi_K^* \} / [\psi_K \psi_K^*], \quad (2)$$

where  $[\psi_K \psi_K^*]$  denotes the average value of  $\psi_K \psi_K^*$  for all points of space. In the special case of free electrons ( $U_K = \text{const.}$ )  $\mathbf{v}_K$  is related to  $\mathbf{K}$  by the equation

$$\mathbf{v}_K = (\hbar/2\pi m) \mathbf{K}$$

† Initiated by F. Bloch, 'Z. Physik,' vol. 52, p. 555 (1928)

For the more general case the relation

$$\mathbf{v}_K = -\frac{2\pi}{h} \text{grad}_K E_K \quad (1)$$

has been suggested, and proved in certain particular cases †. One purpose of this paper is to give a general proof (§ 3).

Secondly, we investigate the effect of uniform electric and magnetic fields upon the Fermi distribution. Kikuchi and Nordheim‡ have shown that if  $f(\mathbf{K}, t)$  is the Fermi distribution function, then if we take an electron with a wave function

$$\psi(x, y, z, t) = \iiint g(\mathbf{K}, t) \psi_{\mathbf{K}}(x, y, z) dK_x dK_y dK_z \quad (2A)$$

which satisfies the initial condition

$$|g(\mathbf{K}, 0)|^2 = f(\mathbf{K}, 0)$$

then for all later times

$$\frac{\partial f}{\partial t} = \frac{\partial}{\partial t} |g|^2$$

In the presence of an electric field  $\mathbf{F}$ ,  $|g(\mathbf{K}, t)|^2$  will be determined by the wave equation

$$\frac{h}{2\pi i} \frac{\partial \psi}{\partial t} = \frac{h^2}{8\pi^2 m} \nabla^2 \psi - V\psi + e\mathbf{F} \cdot \mathbf{R}\psi \quad (3)$$

In the case of free electrons it is easy to show that  $\mathbf{F}$  produces a translation of constant velocity  $(2\pi/h)e\mathbf{F}$ , that is

$$\frac{\partial}{\partial t} |g|^2 = -\frac{2\pi}{h} e\mathbf{F} \cdot \text{grad}_K |g|^2 \quad (11)$$

and various attempts§ have been made to give a general proof. In § 4 we show that (11) is a direct consequence of (3), provided the excited bands (§ 2)

† Bloch, *loc cit*, gave a proof of I in the limiting case of tightly bound electrons.

Pearls, 'Z. Physik,' vol. 53, p. 255 (1929), gave a general proof with the assumption that  $U_K$  varies very slowly with  $\mathbf{K}$ .

‡ 'Z. Physik,' vol. 60, p. 652 (1930).

§ Bloch, *loc cit*, gave a proof where  $|g(\mathbf{K})|$  has the form of a Gauss error function, and in which the excited bands were neglected.

Pearls, *loc cit*, attempted a general proof. The unsatisfactory points in his proof are specifically noted in § 4.

Pearls, 'Ergebn. Exak. Naturwiss.,' vol. 11, p. 279 (1932), has pointed out that the principle of conservation of energy, combined with I, requires that the average of a wave packet in  $\mathbf{K}$  space have the velocity  $(2\pi/h)e\mathbf{F}$ .

are ignored. Further, we shall show that when  $F$  is so small that the probability that the electron makes transitions from one band to another is negligible,† then (II) is true, provided perturbed  $\psi_K$ 's are taken in (2A) which bear the same relation to the unperturbed wave functions as the Stark effect functions do to the unperturbed functions of an atom.

The energy and velocity of these perturbed wave functions are still related by (I)

In the presence of a magnetic field no velocity can be associated with a stationary state  $\psi_K$ , in analogy with there being no unique relation between velocity and momentum in classical mechanics. Nevertheless, if a wave packet be represented by

$$\psi(x, y, z, t) = e^{iA(\mathbf{R}, t)} \iiint g(\mathbf{K}, t) \psi_K dK_x dK_y dK_z \quad (4)$$

the gauge invariance‡ of the wave equation may be so used that the average velocity  $[v]$  of the wave packet is

$$\frac{2\pi}{h} \iiint (\text{grad}_K E_K) |g|^2 dK_x dK_y dK_z \quad (5)$$

We then find (§ 5) that  $|g(\mathbf{K}, t)|^2$  satisfies the equation

$$\frac{\partial}{\partial t} |g|^2 = \frac{\pi e}{h} \mathbf{H} \cdot (\mathbf{v}_K + [v]) \times \text{grad}_K |g|^2 \quad (\text{III})$$

for an interval of time short in comparison to the period of the Larmor precession. Thus when  $|g|^2$  is such a sharp function of  $\mathbf{K}$  that  $[v]$  becomes identical with the velocity  $\mathbf{v}_{K(0)}$  at the centre of the wave packet, the wave packet  $|g|^2$  moves along a surface of constant energy with the velocity  $(2\pi e/hc) \mathbf{v}_{K(0)} \times \mathbf{H}$ .

## § 2 Properties of the Eigenfunctions

The properties of  $\psi_K$  essential for the analysis of the following paragraphs may be derived from a general consideration of the differential equation (1). It will be sufficient to analyse the one dimensional wave equation

$$\left\{ \frac{h^2}{8\pi^2 m} \frac{d^2}{dx^2} - V + E_K \right\} \psi_K = 0 \quad (6)$$

with

$$V(x + na) = V(x) \quad (7)$$

† Zener, communicated to the Royal Society.

‡ Pauli, 'Handbuch d. Physik,' 2nd ed., vol. 24/1, p. 110 (1933).

It may be shown by a variety of methods† that the solutions which remain everywhere finite have the form

$$\psi_k = e^{ikx} U_k(x), \quad (8A)$$

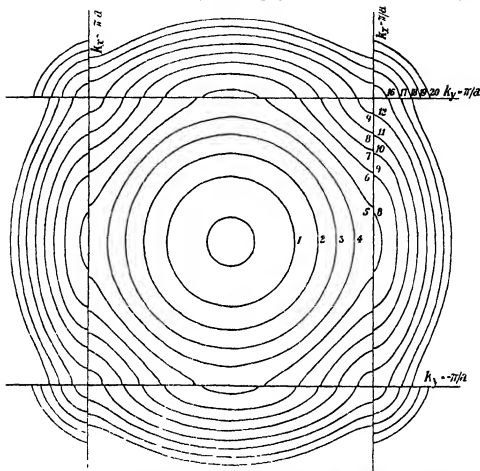


FIG. 1—The curves represent the contours of the energy in the plane  $k_x, k_y$  for the special case when the potential is  $V_0(\cos 2\pi x/a + \cos 2\pi y/a)$  with  $V_0 = 3$  electron volts and  $a = 2.5 \text{ \AA}$ . The numbers by the contour lines give the energy in electron volts.

where  $U_k$  has the same periodic properties as  $V, \pi \epsilon$ ,

$$U_k(x + na) = U_k(x) \quad (8B)$$

† (1) Floquet's theory, Whittaker and Watson, "Modern Analysis," p. 412. (2) Group theory, Bloch, *loc. cit.* (3) An expansion of  $\psi_k$  in a series of exponential functions shows that only those terms combine which differ by the factor  $\exp(2\pi i n x/a)$ . Any sum of such terms will thus have the form (8).

Since  $\psi_k$ ,  $\psi_k^*$ ,  $\psi_{-k}$ , have the same eigenvalues  $E_k$ , a linear relation must exist between these three functions. Inspection shows that this relation can only be

$$\psi_k^* = \psi_{-k} \quad (9)$$

The case in which

$$V = V_0 \cos(2\pi x/a) \quad (10)$$

$V_0$  being a constant, has received special investigation. One result† of particular importance to us is that two distinct solutions having the periodicity  $2a$  cannot have the same eigenvalue. Thus if  $\psi_{n\pi/a}$  and  $\psi_{-n\pi/a}$  have the same eigenvalues, this theorem implies that

$$\psi_{n\pi/a} = \psi_{-n\pi/a} \quad (11)$$

Moreover, the energy as a function of  $k$  has discontinuities at  $k = n\pi/a$ , the two solutions on each edge of the energy discontinuity having an equal claim to the notation  $\psi_k$ .

By employing the double suffix  $(N, K)$  we shall avoid this ambiguity in notation. The label  $N$  may assume the values

$$N = 0, 1, 2, \dots,$$

and  $K$  has the range

$$\pi/a < K \leq \pi/a \quad (12)$$

The relation between the old suffix  $k$  and the new suffixes  $N, K$  is given by

$$k = K \pm \frac{2\pi}{a} N, \quad k \geq 0 \quad (13)$$

These relations do not specify  $N$  and  $K$  uniquely when  $k = n\pi/a$ . We need the additional condition that  $\psi_{N, K}$  is to be a continuous function of  $K$  in the closed interval (12).

Since  $\exp(2\pi i N x/a)$  has the periodicity of the lattice it will be convenient to include this factor in  $U_k$ . We shall thus write

$$\psi_{N, K} = e^{i k x} U_{N, K}$$

where

$$U_{N, K} = e^{\pm 2\pi i N x/a} U_k, \quad K \leq 0$$

Equation (11) now becomes,

$$\psi_{N, -K} = \psi_{N, K} \quad (14)$$

We may thus regard  $K$  as a cyclic variable, the end points of the interval (12) being equivalent.

† Ince, "Ordinary Differential Equations," p. 177, Longmans, Green & Co. (1927)



The convenience of this notation is illustrated by the perspective energy diagram in fig 2

It is of interest to note that equation (9), together with (11), show that  $\psi_{n,n}(x)$ , and  $\psi_{n,n/2}(x) = \psi_{n,-n/2}(x)$  are real functions and hence are associated with a zero average velocity

In the above analysis of the eigenfunctions of (6) we have assumed the special potential (10) The authors are aware of no rigorous proof† for a more general potential The following argument, however, indicates that

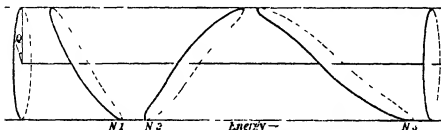


FIG 2—This figure shows diagrammatically the dependence of energy upon  $N$  and  $K$   
 $Q = \alpha K$

the conclusions embodied in equations (12), (13), and (14), are of general validity

Any periodic potential having a physical significance may be represented by the sum

$$V(x) = \sum_{n=1}^M C_n \cos 2\pi n x / a,$$

where  $M$  is a finite number Let  $\psi_{n,k}^{(0)}$  be the eigenfunctions when all the  $C$ 's are zero except  $C_1$  Keep  $C_1$  fixed and let all the other  $C$ 's have finite but arbitrarily small values Let the new eigenfunctions be  $\psi_{n,k}^{(1)}$  These new functions may be expressed by means of the perturbation theory in terms of the  $\psi_{n,k}^{(0)}$ 's Since the expansion will be uniformly convergent, the  $\psi_{n,k}^{(1)}$ 's will be continuous functions of  $k$  in the interval (12), and

† A general proof has been given by M. J. O. Strutt, "Lamé'sche-Mathieu'sche- und Verwandte Funktionen in Physik und Technik," pp 22-23, Springer (1932) His proof however, implicitly assumes that the indefinite integral  $\int (\psi_{n,n/2})^{-2} dx$  has not the period of  $2a$  This assumption needs to be justified, particularly since Strutt's method shows that the indefinite integral  $\int (\psi_{n,n/2})^{-2} dx$ , with  $n > 1$ , must have the period  $2na$  in order that  $\psi_{n,n/2}$  and its complex conjugate be two distinct solutions, each with period  $2na$

will, of course, satisfy (14). They will thus have the same properties as the  $\psi^{(0)}_{\mathbf{k}}$ 's. As we now continuously increase the coefficients, it appears improbable that the  $\psi^{(1)}_{\mathbf{k}}$ 's will not remain continuous functions of  $\mathbf{k}$ . The conclusions obtained for the special potential (10) are thus valid for a general periodic potential, with  $C_1 \neq 0$ .

This analysis may at once be applied to the three dimensional wave equation (3), provided  $V$  is separable in the form  $V = V(x) + V(y) + V(z)$ . The solutions will be labelled by  $\mathbf{N}$  and  $\mathbf{K}$ , where  $\mathbf{N}$  stands for  $(l, m, n)$ ,  $l, m, n$  assume all positive integral values, and where  $\mathbf{K}$  stands for the vector  $(K_x, K_y, K_z)$ , with  $-\pi/a < K_x, K_y, K_z < \pi/a$ . Moreover  $\psi_{\mathbf{K}}$  is a continuous function of  $\mathbf{K}$  such that

$$\psi_{\mathbf{K}} = e^{i\mathbf{K} \cdot \mathbf{R}} U_{\mathbf{K}}(x, y, z), \quad (15)$$

where  $U_{\mathbf{K}}(x, y, z)$  has the same periodicity as  $V(x, y, z)$  and

$$\left. \begin{aligned} \psi_{\mathbf{K} - \pi/a \mathbf{h}_y \mathbf{h}_z} &= \psi_{\mathbf{K} + \pi/a \mathbf{h}_y \mathbf{h}_z} \\ \psi_{\mathbf{K} - \pi/a \mathbf{h}_x \mathbf{h}_z} &= \psi_{\mathbf{K} + \pi/a \mathbf{h}_x \mathbf{h}_z} \\ \psi_{\mathbf{K} - \pi/a \mathbf{h}_x \mathbf{h}_y} &= \psi_{\mathbf{K} + \pi/a \mathbf{h}_x \mathbf{h}_y} \end{aligned} \right\} \quad (16)$$

Brillouin† has shown that a similar grouping occurs when  $V$  may not be so separated. The construction of the closed intervals from the initial  $\mathbf{K}$  space becomes, however, more complicated.

### § 3 Proof of (1)

The second subscript  $\mathbf{N}$  need not be introduced in this paragraph.

In what follows we shall use the vector operators  $\nabla_{\mathbf{K}}, \nabla_{\mathbf{R}}$  defined as follows

$$\begin{aligned} \nabla_{\mathbf{K}} &= \partial/\partial K_x, \partial/\partial K_y, \partial/\partial K_z \\ \nabla_{\mathbf{R}} &= \partial/\partial x, \partial/\partial y, \partial/\partial z \end{aligned}$$

We shall also use the operator  $L_{\mathbf{K}}$  defined by

$$L_{\mathbf{K}} = \frac{\hbar^2}{8\pi^2 m} \nabla^2 + E_{\mathbf{K}} - V$$

Further, the volume integral over a unit cell will be denoted by  $\int_{\text{cell}} d\tau$ .

In this notation the wave equation (1) for the eigenfunctions becomes

$$L_{\mathbf{K}} \psi_{\mathbf{K}} = 0 \quad (17)$$

† "Quantenstatistik," p. 287, Springer (1931)

Since this equation is true for all  $\mathbf{K}$ , we can operate on it with  $\nabla_{\mathbf{K}}$ . Remembering that  $\psi_{\mathbf{K}} = e^{i\mathbf{K} \cdot \mathbf{R}} U_{\mathbf{K}}$ , we obtain

$$(\nabla_{\mathbf{K}} E_{\mathbf{K}}) \psi_{\mathbf{K}} + i L_{\mathbf{K}} \mathbf{R} \psi_{\mathbf{K}} + L_{\mathbf{K}} [e^{i\mathbf{K} \cdot \mathbf{R}} \nabla_{\mathbf{K}} U_{\mathbf{K}}] = 0 \quad (18)$$

It is clear that

$$L_{\mathbf{K}} \mathbf{R} \psi_{\mathbf{K}} = (\hbar^2/4\pi^2 m) \nabla_{\mathbf{K}} \psi_{\mathbf{K}}$$

Hence (18) becomes

$$(\nabla_{\mathbf{K}} E_{\mathbf{K}}) \psi_{\mathbf{K}} + (\hbar^2/4\pi^2 m) i \nabla_{\mathbf{K}} \psi_{\mathbf{K}} + L_{\mathbf{K}} [e^{i\mathbf{K} \cdot \mathbf{R}} \nabla_{\mathbf{K}} U_{\mathbf{K}}] = 0 \quad (19)$$

If we multiply (19) by  $\psi_{\mathbf{K}}^*$ , and integrate over a unit cell, we obtain

$$(\nabla_{\mathbf{K}} E_{\mathbf{K}}) \int_c \psi_{\mathbf{K}}^* \psi_{\mathbf{K}} d\tau + \frac{i\hbar^2}{4\pi^2 m} \int_c \psi_{\mathbf{K}}^* \nabla_{\mathbf{K}} \psi_{\mathbf{K}} d\tau + \int_c \psi_{\mathbf{K}}^* L_{\mathbf{K}} [e^{i\mathbf{K} \cdot \mathbf{R}} \nabla_{\mathbf{K}} U_{\mathbf{K}}] d\tau = 0$$

The last term may be shown to vanish identically, for since the integrand has the same periodicity as the lattice, we obtain, on integrating twice by parts with respect to  $x, y, z$ ,

$$\int_c \psi_{\mathbf{K}}^* L_{\mathbf{K}} [e^{i\mathbf{K} \cdot \mathbf{R}} \nabla_{\mathbf{K}} U_{\mathbf{K}}] d\tau = \int_c [e^{i\mathbf{K} \cdot \mathbf{R}} \nabla_{\mathbf{K}} U_{\mathbf{K}}] L_{\mathbf{K}} \psi_{\mathbf{K}}^* d\tau$$

But since  $L_{\mathbf{K}}$  is a real operator  $L_{\mathbf{K}} \psi_{\mathbf{K}}^*$  vanishes identically by (17)

We now have

$$\frac{\hbar}{2\pi i m} \int_c \psi_{\mathbf{K}}^* \nabla_{\mathbf{K}} \psi_{\mathbf{K}} d\tau \Big/ \int_c \psi_{\mathbf{K}}^* \psi_{\mathbf{K}} d\tau = \frac{2\pi}{\hbar} \nabla_{\mathbf{K}} E_{\mathbf{K}}$$

But the left member of this equation is equivalent to the definition (2) of  $v_{\mathbf{K}}$ . For integration by parts gives

$$\int_c \psi_{\mathbf{K}}^* \nabla_{\mathbf{K}} \psi_{\mathbf{K}} d\tau = - \int_c \psi_{\mathbf{K}} \nabla_{\mathbf{K}} \psi_{\mathbf{K}}^* d\tau$$

Hence

$$\begin{aligned} \int_c \psi_{\mathbf{K}}^* \nabla_{\mathbf{K}} \psi_{\mathbf{K}} d\tau &= \frac{1}{2} \int_c \{ \psi_{\mathbf{K}}^* \nabla_{\mathbf{K}} \psi_{\mathbf{K}} - \psi_{\mathbf{K}} \nabla_{\mathbf{K}} \psi_{\mathbf{K}}^* \} d\tau \\ &= \frac{1}{2} V_0 \{ \psi_{\mathbf{K}}^* \nabla_{\mathbf{K}} \psi_{\mathbf{K}} - \psi_{\mathbf{K}} \nabla_{\mathbf{K}} \psi_{\mathbf{K}}^* \} \end{aligned}$$

where  $V_0$  is the volume of the unit cell

We have thus established relation (I)

#### § 4 Proof of (II)

In this section we prove relation (II) directly from the wave equation (3). Using the notation introduced in § 2, we may expand the solution of this equation as†

$$\psi(x, y, z, t) = \sum_N \iiint_{-\pi/a}^{\pi/a} g_N(\mathbf{K}, t) \psi_{\mathbf{K}}(x, y, z) dK_x dK_y dK_z \quad (20)$$

† *Pearls, loc. cit.*, represented the wave packet by a summation over the discrete set of quantum numbers obtained by specifying periodic boundary conditions. This summation

The  $\psi_{N, \mathbf{k}}$  are the eigenfunctions of the unperturbed equation (1). They will be taken to be normalized to a unit cell, so that

$$\int_{\tau} \psi_{N, \mathbf{k}}^* \psi_{N, \mathbf{k}} d\tau = \delta_{N, N'} \quad (21)$$

where, as in the previous section,

$$\int_{\tau} d\tau \text{ denotes } \iiint_{\Omega} dx dy dz$$

In order to obtain the wave equation for the  $q_{N, \mathbf{k}}(K, t)$ , we substitute the expansion (20) into equation (3), multiply by  $\psi_{N, \mathbf{k}}^*$ , and integrate over all co-ordinate space. The integrals are evaluated in the appendix. Denoting by  $H''$  the operator

$$H'' = H_0 - eF (\mathbf{R} + i\nabla_{\mathbf{K}}), \quad (22)$$

where  $H_0$  is the Hamiltonian for the unperturbed lattice, and writing

$$(N|H''|N')_{\mathbf{k}} = \int_{\tau} \psi_{N, \mathbf{k}}^* H'' \psi_{N', \mathbf{k}} d\tau,$$

we obtain

$$\left( \frac{\hbar}{2\pi} \frac{\partial}{\partial t} + eF \cdot \nabla_{\mathbf{K}} \right) q_{N, \mathbf{k}}(K, t) = - \sum_{N'} (N|H''|N')_{\mathbf{k}} q_{N'}(K, t) \quad (23)$$

For further reduction we must use the Hermitian property of the operator  $H''$ . The Hermitian property of all the components of  $H''$  except  $eF \cdot i\nabla_{\mathbf{K}}$  is obvious. The property for  $i\nabla_{\mathbf{K}}$  is established by operating upon equation (21) with  $i\nabla_{\mathbf{K}}$ . We obtain at once the desired relation

$$(N|i\nabla_{\mathbf{K}}|N')_{\mathbf{k}} = (N'|i\nabla_{\mathbf{K}}|N)_{\mathbf{k}}^*$$

We now multiply (23) by  $g_{N, \mathbf{k}}^*(K, t)$ , and add the resulting equation to its conjugate complex. We obtain

$$\left( \frac{\hbar}{2\pi} \frac{\partial}{\partial t} + eF \cdot \nabla_{\mathbf{K}} \right) |g_N|^2 = \sum_N \{ (N|H''|N')_{\mathbf{k}} g_{N'}^* g_N - (N'|H''|N)_{\mathbf{k}} g_N^* g_N \}$$

is intrinsically more difficult to handle than an integral. Peierls completed his analysis by using the following approximations. In the sum

$$\sum_{N'}^{G/2-1} \alpha \exp 2\pi i (k - k') \alpha / G = G \cos \pi (k - k') (e^{2\pi i (k - k') / G} - 1)^{-1}$$

he replaced the oscillating numerator,  $\cos \pi (k - k')$ , by unity. In calculating the sum

$$\sum_L C(k) (e^{2\pi i (k - k') / G} - 1)^{-1}$$

he assumes that  $C(k)$  is very small except when  $|k - k'| \ll G$ , which is certainly not true of the Fermi function in actual metals.

Since the bracketed expression in the right member is antisymmetric with respect to an interchange of  $N$  and  $N'$ , the right member vanishes when we sum over  $N$ . We are left with

$$\left( \frac{\hbar}{2\pi} \frac{\partial}{\partial t} + eF \cdot \nabla_{\mathbf{K}} \right) \sum_{\mathbf{N}} |g_{\mathbf{N}}|^2 = 0 \quad (24)$$

The solution of this equation is, of course,

$$\sum_{\mathbf{N}} |g_{\mathbf{N}}(\mathbf{K}, t)|^2 = G \left( \frac{\hbar \mathbf{K}}{2\pi} - eFt \right)$$

where  $G$  is an arbitrary function of its argument. The function  $\sum_{\mathbf{N}} |g_{\mathbf{N}}(\mathbf{K}, t)|^2$  thus moves with the uniform velocity  $2\pi eF/\hbar$ .

Suppose now that *initially* the electron is definitely in one band so that all the  $g_{\mathbf{N}}(\mathbf{K})$  vanish except, say,  $g_1(\mathbf{K})$ . Then if we assume that also after time  $t$  the other bands may be left out of account,† only  $g_1$  will remain in the summation of (24), and we have immediately relation (11).

The upper bands affect the problem in two different ways. The wave function  $\psi_{\mathbf{N}, \mathbf{K}}$  will become perturbed *as soon as* the field is put on, in the same way that an atomic function is perturbed by an electric field. We may reasonably expect that the effect will be given by a first order perturbation calculation, as for a free atom. The wave function may be expanded in terms of all the  $\psi_{\mathbf{N}, \mathbf{K}}$ , but this does *not* mean that the electron actually gets into the excited band. On the other hand, there is a probability, which increases with time, that the electron will make a transition from one band to the next. This transition has been discussed by Zener (*loc cit*) and is always negligible in metals.

Consider the equation

$$H'' \psi''_{\mathbf{N}, \mathbf{K}} = E''_{\mathbf{N}, \mathbf{K}} \psi''_{\mathbf{N}, \mathbf{K}},$$

where, as in (22),

$$H'' = H_0 - eF(\mathbf{R} + i\nabla_{\mathbf{K}})$$

Let  $\psi''_{\mathbf{N}, \mathbf{K}}$  be the solution of this equation which is obtained when  $eF(\mathbf{R} + i\nabla_{\mathbf{K}})$  is treated as a small perturbation. The first order perturbation calculation‡ gives

$$\psi''_{\mathbf{N}, \mathbf{K}} = \psi_{\mathbf{N}, \mathbf{K}} + \sum_{\mathbf{N}'} \frac{(N | eF(\mathbf{R} + i\nabla_{\mathbf{K}}) | N')_{\mathbf{K}}}{E_{\mathbf{N}, \mathbf{K}} - E_{\mathbf{N}', \mathbf{K}}} \psi_{\mathbf{N}', \mathbf{K}}$$

† This was tacitly assumed in the restricted proof of Bloch, *loc cit*, and in the attempted general proof of Peierls, *loc cit*.

‡ The higher order approximations will not be given by the customary formulae, since the operator  $\nabla_{\mathbf{K}}$  in  $H''$  does not commute with the expansion coefficients.

or, omitting the common factor  $e^{i\mathbf{K} \cdot \mathbf{R}}$ ,

$$U''_{\mathbf{N}, \mathbf{K}} = U_{\mathbf{N}, \mathbf{K}} + \sum_{\mathbf{N}'} \frac{(\mathbf{N} | e\mathbf{F} \cdot (\mathbf{R} + \mathbf{r}) | \mathbf{N}')_{\mathbf{K}}}{E_{\mathbf{N}, \mathbf{K}} - E_{\mathbf{N}', \mathbf{K}}} U_{\mathbf{N}', \mathbf{K}}$$

These approximate functions will not form exactly a complete orthogonal set. Nevertheless, if the electron is initially in the  $N$ th band, we may expect that  $\psi$  as represented by the expansion

$$\psi(x, y, z, t) = \iiint_{\pi a}^{\pi a} g''_{\mathbf{N}}(\mathbf{K}, t) \psi''_{\mathbf{N}, \mathbf{K}}(x, y, z) dK_x dK_y dK_z$$

will differ inappreciably from the exact  $\psi$  until transitions to the adjacent bands become important.

We now substitute this expansion into the wave equation (3), multiply by  $\psi''^*_{\mathbf{N}, \mathbf{K}}$ , and integrate over all co-ordinate space. Using the orthogonality relation

$$\int \psi''_{\mathbf{N}, \mathbf{K}} \psi''^*_{\mathbf{N}', \mathbf{K}} d\tau = \delta_{\mathbf{N}, \mathbf{N}'}$$

and the integrals in the appendix, we obtain

$$\left( \frac{\hbar}{2\pi} \frac{\partial}{\partial t} + e\mathbf{F} \cdot \nabla_{\mathbf{K}} \right) g''_{\mathbf{N}}(\mathbf{K}, t) = iE''_{\mathbf{N}, \mathbf{K}} g''_{\mathbf{N}}(\mathbf{K}, t)$$

We thus obtain

$$\left( \frac{\hbar}{2\pi} \frac{\partial}{\partial t} + e\mathbf{F} \cdot \nabla_{\mathbf{K}} \right) |g''_{\mathbf{N}}(\mathbf{K}, t)|^2 = 0$$

and so

$$|g''_{\mathbf{N}}(\mathbf{K}, t)|^2 = \text{const} \left( \frac{\hbar}{2\pi} \mathbf{K} - e\mathbf{F}t \right)$$

Neglecting transitions between bands, the action of the electric field has thus two distinct effects. Firstly, it polarizes the individual "atomic" functions  $U_{\mathbf{N}, \mathbf{K}}$ . Secondly, it produces a uniform translation of  $|g''_{\mathbf{N}}|^2$  in  $\mathbf{K}$  space.

It is of interest to note that the expression for the polarized "atomic" functions becomes, when the lattice constant is increased indefinitely, identical to the usual expression for the polarized wave functions of isolated atoms. For large values of the lattice constant  $\psi_{\mathbf{N}, \mathbf{K}}$  becomes independent of  $\mathbf{K}$  apart from a constant factor, in the region where  $\psi_{\mathbf{N}, \mathbf{K}}$  is non-vanishing. The second term in the perturbation thus becomes negligible.

It is also of interest to note that the velocity  $v''_{N, K}$  associated with a polarized wave function  $\psi''_{N, K}$  is related to the energy  $E''_{N, K}$  by a relation analogous to (I), that is

$$v''_{N, K} = \frac{2\pi}{h} \text{grad}_K E''_{N, K}$$

When we utilize the Hermitian property of the operator  $H''$  the proof follows that given in § 3 for equation (I).

### § 5 Investigation of (111)

In this section we investigate the motion of a wave packet in a lattice under the influence of a magnetic field  $H$ . Since the equations which we obtain are valid only for short intervals of time, it would not be profitable to introduce the excited states. Hence in place of representing the wave packet by the sum

$$\psi(x, y, z, t) = \sum_N \iiint_{-\pi/a}^{+\pi/a} g_N(K) \psi_{N, K} dK_x dK_y dK_z,$$

which we would do, of course, if we were to be exact, we shall take only one term of this sum, and so shall omit the suffix  $N$ . Hence the wave packet will be represented by

$$\psi(x, y, z, t) = \iiint_{-\pi/a}^{+\pi/a} g(K) \psi_K dK_x dK_y dK_z, \quad (25)$$

We shall take  $\psi$  to be normalized, i.e.,

$$\int \psi^* \psi d\tau = 1$$

The function  $\psi$  must satisfy the wave equation

$$\left\{ \frac{1}{2m} \left( \frac{h}{2\pi i} \nabla - \frac{e}{c} \mathbf{A} \right)^2 + V + \frac{h}{2\pi i} \frac{\partial}{\partial t} \right\} \psi(x, y, z, t) = 0 \quad (26)$$

The only condition which the vector potential  $\mathbf{A}$  must satisfy is

$$\text{curl } \mathbf{A} = \mathbf{H}$$

The general solution of this equation may be written as

$$\mathbf{A} = \frac{1}{2} \mathbf{H} \times \mathbf{R} + \nabla f,$$

where  $f$  is an arbitrary function of  $\mathbf{R}$  and of  $t$ . The scalar potential  $V$  is then given by

$$V = V_0 - \frac{e}{c} \frac{\partial f}{\partial t},$$

where  $V_0$  is the periodic potential due to the lattice

The arbitrariness of  $f$  will be used to simplify the expression for the average velocity  $[v]$  of the wave packet. This average velocity is given by

$$[v] = \frac{\hbar}{4\pi m_1} \int \{\psi^* \nabla \psi - \psi \nabla \psi^*\} d\tau - \frac{e}{mc} \int \mathbf{A} \psi \psi^* d\tau$$

Now if we choose

$$\mathbf{A} = \frac{1}{2} \mathbf{H} \times (\mathbf{R} - [\mathbf{R}]), \quad (27)$$

where

$$[\mathbf{R}] = \int \psi^* \mathbf{R} \psi d\tau,$$

then clearly

$$\int \mathbf{A} \psi \psi^* d\tau = 0,$$

and the average velocity becomes

$$[v] = \iiint_{-\pi/a}^{\pi/a} \mathbf{v}_K |g(\mathbf{K})|^2 dK_x dK_y dK_z,$$

where  $\mathbf{v}_K$  is defined by (2). In order to write  $\mathbf{A}$  in this form, we must take

$$f = -\frac{1}{2} \mathbf{R} \cdot \mathbf{H} \times [\mathbf{R}] = \frac{1}{2} \mathbf{H} \cdot \mathbf{R} \times [\mathbf{R}]$$

Then since

$$\frac{\partial}{\partial t} [\mathbf{R}] = [v],$$

the scalar potential becomes

$$V = V_0 - \frac{e}{2c} \mathbf{H} \cdot \mathbf{R} \times [v] \quad (28)$$

Taking the vector and scalar potentials to be given by (27) and (28), we substitute the expansion (25) into (26), multiply by  $\psi_K^*$ , and integrate over all co-ordinate space. We obtain, using (A6), (A7),

$$\begin{aligned} \frac{\partial}{\partial t} g(\mathbf{K}) &= -\frac{2\pi v}{h} E_K g(\mathbf{K}) + \frac{2\pi v}{h} \left(\frac{a}{2\pi}\right)^3 \int \left( \left| \int_{-\pi/a}^{\pi/a} g(\mathbf{K}') \psi_K^* \right. \right. \\ &\quad \times \left. \left\{ \frac{\hbar}{2\pi v} \frac{e}{2mc} \mathbf{H} \cdot (\mathbf{R} - [\mathbf{R}]) \times \nabla + \frac{e}{2c} \mathbf{H} \cdot \mathbf{R} \times [v] \right\} \right. \\ &\quad \left. \left. - \frac{e^2}{8mc^2} [\mathbf{H} \times (\mathbf{R} - [\mathbf{R}])]^2 \right\} \psi_K dK'_x dK'_y dK'_z \right) d\tau. \end{aligned} \quad (29)$$

We have, of course, utilized the fact that the  $\psi_K$  are eigenfunctions of

$$\{(\hbar^2/8\pi^2 m_1) \nabla^2 + E_K - V_0\} \psi_K = 0$$



We multiply equation (29) by  $g^*(\mathbf{K})$ , and add the resulting equation to its conjugate complex. The integrals are evaluated by the method outlined in the appendix. Neglecting terms in  $H^2$ , we obtain

$$\frac{\partial}{\partial t} |g|^2 = \frac{2\pi e}{h} \frac{1}{2c} \mathbf{H} \cdot (\mathbf{v}_\mathbf{K} + [\mathbf{v}]) \times \nabla_\mathbf{K} |g|^2 \quad (30)$$

We now consider the terms which have been neglected in (30). In their evaluation, we need to know the phase of  $g$ . If we neglect powers of  $H$  greater than the second, we may take

$$g = |g| e^{2\pi i \mathbf{R} \cdot \mathbf{K} / h}$$

We separate the terms in  $H^2$  into those that are also present for free electrons, and those that vanish for free electrons. The former may be written as

$$\frac{2\pi e^2}{h} \frac{1}{4mc^2} (\mathbf{H} \times \nabla_\mathbf{K}) \cdot \{\mathbf{H} \times (\mathbf{v}_\mathbf{K} t - [\mathbf{R}])\} |g|^2$$

If  $|g(\mathbf{K})|^2$  is a sufficiently sharp function of  $\mathbf{K}$ , the variation of  $\mathbf{v}_\mathbf{K}$  may be neglected in comparison with the variation of  $|g|^2$ . This term then becomes

$$\frac{2\pi e^2}{h} \frac{1}{4mc^2} \{\mathbf{H} \times (\mathbf{v}_\mathbf{K} t - [\mathbf{R}])\} \cdot \{\mathbf{H} \times \nabla_\mathbf{K} |g|^2\} \quad (31)$$

The origin of co-ordinates may be so chosen that  $[\mathbf{R}] = 0$  at  $t = 0$ . We then observe that the ratio of (31) to the right member of (30) is approximately  $(eH/2mc)\omega t$ , i.e.,  $\omega t$ , where  $\omega$  is the Larmor precession frequency. Hence (31) may be neglected only for an interval of time  $t$  such that  $\omega t \ll 1$ .

Those terms in  $H^2$  which vanish for free electrons reduce to

$$\frac{2\pi}{h} \left(\frac{2\pi}{a}\right)^3 \frac{e^2}{4\pi u^2} \{\mathbf{H} \times \int_e \mathbf{U}^*_{\mathbf{K}'} \nabla_\mathbf{K} \mathbf{U}_\mathbf{K} d\tau\} \cdot \{\mathbf{H} \times \nabla_\mathbf{K} |g|^2\}$$

This may, of course, always be neglected in comparison to the right member of (30) when  $H$  is sufficiently small.

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### Summary

In the modern literature of the theory of metallic conduction certain equations which are true for free electrons are assumed to remain valid when the potential

is periodic. At best proofs have been given only for particular cases. In this paper general proofs are given for these equations. These equations give (I) the relationship between the velocity and energy of a stationary state, (II) the effect of a constant electric field upon a wave packet, (III) the effect of a constant magnetic field upon a wave packet.

## APPENDIX

Let  $\phi(k, x)$  be a continuous function of its arguments having the periodic property

$$\phi(k, x + a) = \phi(k, x)$$

Then

$$\begin{aligned} J &\equiv \int_{-\infty}^{\infty} \left\{ \int_{k_1}^{k_2} e^{i(k-k')x} \phi(k, x) dk \right\} dx \\ &= \frac{2\pi}{a} \int_0^a \phi(k', x) dx, \quad k_1 < k' < k_2 \end{aligned} \quad (A1)$$

When  $\phi(k, x)$  is independent of  $x$ , this integral reduces to the Fourier Integral. The proof of (A1) which we give below in fact follows closely the ordinary proof† of the Fourier Integral.

The integral

$$\int_{k_1}^{k_2} e^{i(k-k')x} \phi(k, x) dk$$

represents a wave packet in  $x$  space, and so vanishes at  $x = \pm \infty$ . Hence we may write

$$J = \lim_{n \rightarrow \infty} \int_{-na}^{na} \int_{k_1}^{k_2} e^{i(k-k')x} \phi(k, x) dk dx$$

The order of these finite integrations may now be interchanged. Now

$$\begin{aligned} \int_{-na}^{na} e^{i(k-k')x} \phi(k, x) dx &= \sum_{s=-n}^{n-1} e^{i(k-k')as} q(k) \\ &= \frac{2i \sin(k-k')na}{e^{i(k-k')a} - 1} q(k), \end{aligned}$$

where

$$q(k) = \int_0^a e^{i(k-k')s} \phi(k, s) ds$$

We thus have

$$J = \lim_{n \rightarrow \infty} \int_{k_1}^{k_2} \frac{2i \sin(k-k')na}{e^{i(k-k')a} - 1} q(k) dk$$

† Courant and Hilbert, "Methoden der Mathematischen Physik," vol. 1, pp. 54-55, Berlin (1934).

If  $k'$  were outside the interval  $k_1 < k < k_2$ , integration by parts would show that  $J = 0$ . Hence  $J$  may be written as

$$J = \lim_{n \rightarrow \infty} \int_{k-\varepsilon}^{k+\varepsilon} \frac{2i \sin(k-k')na}{e^{i(k-k')a} - 1} q(k) dk$$

where  $\varepsilon$  is arbitrarily small. We now make  $\varepsilon$  so small that  $q(k)$  may be replaced by  $q(k')$ , and  $\exp i(k-k')a$  by  $1 + i(k-k')a$ .

We thus have finally

$$J = (2/a) q(k') \lim_{n \rightarrow \infty} \int_{-\varepsilon}^{\varepsilon} \frac{\sin n\xi}{\xi} d\xi = \frac{2\pi}{a} \phi(k', x) dx$$

The generalization of (A1) to three dimensions gives

$$\int \left\{ \int e^{i(\mathbf{K}-\mathbf{K}') \cdot \mathbf{R}} \phi(\mathbf{K}, \mathbf{R}) d\mathbf{K} \right\} d\tau = \left( \frac{2\pi}{a} \right)^3 \int_{\varepsilon} \phi(\mathbf{K}', \mathbf{R}) d\tau, \quad (\text{A2})$$

where

$$\int d\mathbf{K} \text{ denotes } \iiint_{-\pi/a}^{\pi/a} dK_x dK_y dK_z,$$

$$\int d\tau \text{ denotes } \iiint_{-\infty}^{\infty} dx dy dz,$$

and

$$\int_{\varepsilon} d\tau$$

represents an integration over a unit cell, and  $\mathbf{K}'$  is restricted to the range  $-\pi/a < K'_x, K'_y, K'_z < \pi/a$ .

The second formula which we need is the orthogonality relation

$$\int_{\varepsilon} \psi_{N, \mathbf{K}}^* \psi_{N', \mathbf{K}} d\tau = 0, \quad N' \neq N \quad (\text{A3})$$

This is proved by the usual procedure of subtracting

$$\psi_{N, \mathbf{K}}^* (H_0 - E_{N, \mathbf{K}}) \psi_{N, \mathbf{K}} = 0$$

from

$$\psi_{N, \mathbf{K}} (H_0 - E_{N', \mathbf{K}}) \psi_{N', \mathbf{K}}^* = 0$$

and integrating over a unit cell. Integration by parts then gives (A3).

In the following applications of these two formulae to the integrals occurring in the text,  $\psi_{N, \mathbf{K}}$  will be taken as normalized over a unit cell.

The simplest of these integrals follows directly

$$\iint h(\mathbf{K}) \psi_{N, \mathbf{K}}^* \psi_{N, \mathbf{K}} d\mathbf{K} d\tau = \delta_{N, N} \left( \frac{2\pi}{a} \right)^3 h(\mathbf{K}') \quad (\text{A4})$$

In the next two integrals we must first replace  $\mathbf{R} \exp(i\mathbf{K} \cdot \mathbf{R})$  by  $-i\nabla_{\mathbf{K}} \exp(i\mathbf{K} \cdot \mathbf{R})$ , and integrate by parts with respect to  $K_x, K_y, K_z$ . Use is made of the cyclic character of  $\mathbf{K}$  (equation (16))

$$\left(\frac{a}{2\pi}\right)^3 \iint g_N(\mathbf{K}) \psi_{\mathbf{K}}^* \mathbf{R} \psi_{\mathbf{K}} d\mathbf{K} d\tau = i\nabla_{\mathbf{K}} g_N(\mathbf{K}') \delta_{\mathbf{K}', \mathbf{N}} + \int_c U_{\mathbf{K}', \mathbf{K}}^* i\nabla_{\mathbf{K}} U_{\mathbf{K}} d\tau g_N(\mathbf{K}') \quad (\text{A5})$$

$$\begin{aligned} \left(\frac{a}{2\pi}\right)^3 \iint g_V(\mathbf{K}) \psi_{\mathbf{K}}^* \mathbf{R} \times \nabla_{\mathbf{R}} \psi_{\mathbf{K}} d\mathbf{K} d\tau \\ \int_c \psi_{\mathbf{K}}^* \nabla_{\mathbf{R}} \psi_{\mathbf{K}} d\tau \times i\nabla_{\mathbf{K}} g_N(\mathbf{K}') \\ - \left\{ \int_c \psi_{\mathbf{K}}^* \nabla_{\mathbf{R}} e^{i\mathbf{K} \cdot \mathbf{R}} \times i\nabla_{\mathbf{K}} U_{\mathbf{K}} d\tau \right\} g_N(\mathbf{K}') \end{aligned} \quad (\text{A6})$$

Finally, in the following integral we first replace  $\mathbf{R}^2 \exp(i\mathbf{K} \cdot \mathbf{R})$  by  $-\nabla_{\mathbf{K}}^2 \exp(i\mathbf{K} \cdot \mathbf{R})$ , and integrate by parts twice with respect to  $K_x, K_y, K_z$

$$\begin{aligned} \left(\frac{a}{2\pi}\right)^3 \iint g_V(\mathbf{K}) \psi_{\mathbf{K}}^* |\mathbf{R}|^2 \psi_{\mathbf{K}} d\mathbf{K} d\tau = -\nabla_{\mathbf{K}}^2 g_V(\mathbf{K}') \\ - 2\{ \nabla_{\mathbf{K}} g_V(\mathbf{K}') \} \int_c U_{\mathbf{K}}^* \nabla_{\mathbf{K}} U_{\mathbf{K}} d\tau \\ - g_N(\mathbf{K}') \int_c U_{\mathbf{K}}^* \nabla_{\mathbf{K}}^2 U_{\mathbf{K}} d\tau \end{aligned} \quad (\text{A7})$$

## *A Photoelectric Spectrophotometer using Dual Electrostatic Compensation*

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(Communicated by N. V. Sidgwick, F.R.S. — Received August 4, 1933)

### I *Introduction*

The photometer here described is suitable for the purpose of measuring extinction coefficients of substances in solution, more particularly in the ultra-violet region. The method employed is a purely electrical one, and does not involve the use of any mechanical devices (sectors or wedges, etc.) for the quantitative variation of light intensity. Errors due to fluctuations of the light source are eliminated by the use of two photoelectric cells, while the simultaneous electrostatic compensation of both cells does away with the necessity of measuring or controlling times of exposure. The actual observations are of (i) a resistance plugged into a resistance box, and (ii) an electrometer deflection. The former gives the approximate result and the latter the correction to be applied.

### II *Description of Apparatus and Method*

The source of light is a vertical mercury vapour lamp, running at about  $3\frac{1}{2}$  amps and about 100 volts terminal voltage. The light from this source is condensed upon the slit of a double quartz monochromator by a suitable cylindrical quartz lens with its axis vertical. By means of the double quartz monochromator,\* which has auxiliary dispersing prisms rotated by means of wave-length drums, any desired wave-length emitted by the source can be selected and allowed to enter the photoelectric photometer.

In the figure, which is purely diagrammatic, the exit slit of the double monochromator is shown at S. After having been collimated by a quartz lens L, the beam of monochromatic light (shown dotted) is incident upon an inclined quartz plate Q, which is partially platinized by sputtering. A fraction of the light is reflected at the plate and enters one of the photoelectric cells  $P_c$ , which we will call the comparison cell. A fraction of the light is transmitted by the plate Q and passes on to the other photoelectric cell  $P_m$ , which we will call the measuring cell. Between Q and the measuring cell can be introduced the

\* Made by Messrs. Bellingham & Stanley, Ltd.

quartz-ended absorption vessel A containing the liquid whose absorption is under investigation

The photoelectric cells are both of the vacuum type. Their cathodes are connected together and maintained at a negative potential by means of a battery  $B_1$ , the positive end of which is earthed. The voltage of  $B_1$  is more

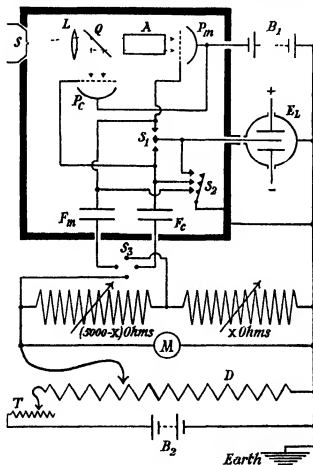


FIG 1

than sufficient to ensure saturation of the cells, so that the photoelectric current is in each case directly proportional to the incident light intensity. The cathode metal is sodium, which gives a sufficient response down to a wavelength of 2500 Å. The envelopes of the cells are made entirely of quartz and the distance between anode and cathode seals ensures high insulation \*

\* Cells specially made by Otto Pressler, of Leipzig.

The rest of the apparatus, to be described and discussed in detail below, provides a means of measuring the ratio of the photoelectric current given by the measuring cell to that given by the comparison cell. Since both cells have a linear response, this ratio is proportional to the ratio of the corresponding incident light intensities. Errors due to fluctuations of the light source are eliminated, for such fluctuations alter the intensities of both beams in the same ratio. The determination of the extinction coefficient of a substance in solution involves the measurement of the ratio of the intensity  $I_0$  incident upon the solution to the intensity  $I$  transmitted by it. When the absorption vessel  $A$  is filled with the solution, the measured ratio of the current given by the measuring cell to that given by the comparison cell is  $kI/I_0$ , where  $I_0$  is the intensity of the beam falling upon the comparison cell and  $k$  is a constant involving the sensitivities of the two cells. This ratio having been determined, the solution is replaced by pure solvent and the corresponding ratio  $kI_0/I_0$  is measured. Division of the second ratio by the first then gives the required quantity  $I_0/I$ . This procedure eliminates the effects due to the absorption vessel and the solvent. It will be noted that the properties of the reflector  $Q$  and the sensitivities of the photoelectric cells do not enter into the final result, provided only that they remain constant over the duration of the determination.

Now as to the method of measuring the ratio of the photoelectric currents. The anodes of the two photoelectric cells are connected respectively to poles of a switch  $S_1$ , by means of which either the one or the other may be put into connection with the needle of the Landemann electrometer  $E_L$ .<sup>\*</sup> The earthing switch  $S_2$  permits of the earthing of both the anodes and the electrometer needle together. In the actual apparatus  $S_1$  and  $S_2$  are constructed as one unit, so designed as to operate without the generation of charges which would disturb the electrometer needle. The contacts are all gold-plated and are gently pressed together without friction. The arrangement for applying the necessary voltages to the plates of the Landemann electrometer has been omitted from the figure for the sake of clarity. The anode of the comparison cell  $P_c$  is further connected to one plate of the screened induction condenser  $F_c$ , while the anode of the measuring cell  $P_m$  is likewise connected to one plate of the similar screened induction condenser  $F_m$ . These are specially constructed cylindrical air condensers† (approximate capacity 16 cm.) An earthed metal casing (shown by a heavy line in the figure) encloses the whole of the apparatus

<sup>\*</sup> F. A. and A. F. Landemann and Keeley, 'Phil. Mag.', vol. 47, p. 578 (1924).

<sup>†</sup> See Campbell and Ritchie, "Photoelectric Cells" (1929), fig. 30, p. 126.

so far described and screens it from external influences. The switches  $S_1$  and  $S_2$  are manipulated by suitable controls projecting through the casing. The short lead to the electrometer needle is screened by an earthed metal tube. The whole of the insulation of the anode systems (switches, induction condensers, etc.) to which the needle may be connected is of amber, with the exception of the photoelectric cells which are of quartz.

The plates of the induction condensers which are not wired to the photoelectric cells are connected through a change-over switch  $S_3$  to the double potentiometer arrangement shown in the lower part of the figure. This consists of a sliding potentiometer  $D$  fed by a battery  $B_1$ , the slider being connected to one end of a second potentiometer formed by two standard variable resistance boxes, marked  $X$  ohms and  $(5000 - X)$  ohms in the figure. The sum of the resistances of the boxes is kept constant (actually equal to 5000 ohms).

The method is as follows. By means of the switch  $S_1$  the electrometer needle is put into connection with one or other of the photoelectric cells, the switch  $S_2$  being closed. Switch  $S_2$  is now opened so as to insulate both cell anodes, and the beam of monochromatic light is switched on by opening a shutter (not shown in the figure). The photoelectric currents given by the two cells begin to charge up their respective anode systems negatively, and the electrometer needle begins to move as a result of the rise of the negative potential of the system of which it forms a part. This needle movement is compensated, i.e., the needle is kept at its original zero position, by moving the slider of the potentiometer  $D$ , which applies a positive potential to the induction condenser concerned and maintains the needle potential at its original zero value by electrostatic induction\*. When the slider of  $D$  has been moved along the whole extent of its traverse, the light is cut off by closing the shutter.

Now during the whole operation positive potentials have been applied to both induction condensers simultaneously, the ratio of these potentials being determined by the ratio of the resistances of the boxes. For a certain value of this ratio, not only will the current from the one photoelectric cell have been exactly compensated, but also the current from the other cell. Under these conditions, therefore, there will be no throw of the electrometer when, after the light has been cut off, the switch  $S_1$  is moved over into its other position. If the resistance ratio has not this correct value, a throw of the electrometer needle will be observed on thus moving over the switch  $S_1$ . Hence we have the possibility of finding by trial the correct value of the resistance ratio giving

\* Cf. Campbell and Ritchie, *loc. cit.*, p. 114.



simultaneous exact compensation of both photoelectric currents. In practice this lengthy process of trial is rendered unnecessary by a method to be described later.

Let  $C_1$  and  $C_2$  be the capacities of the induction condensers connected to the photoelectric cells giving the greater current and the lesser current respectively. If the total voltage applied from the potentiometer  $D$  is  $V$ , the magnitude of the charge on the first condenser will be  $C_1 V$  and that on the second condenser  $C_2 VX/5000$ , where  $X/5000$  is the value of the ratio of the resistance of the one box to the total resistance of both boxes, as indicated in the figure. These charges are equal to the time integrals of the respective photoelectric currents over the duration  $t$  of the operation. Hence when dual compensation is effected,

$$-C_1 V = i_1 t \quad \text{and} \quad -C_2 VX/5000 = i_2 t,$$

where  $i_1$  and  $i_2$  are the respective average photoelectric currents. Hence

$$\frac{i_1}{i_2} = \frac{C_1}{C_2} \frac{5000}{X}$$

The capacity ratio  $C_1/C_2$  is a constant, and hence the determination of  $X$  gives a measure of the ratio of the photoelectric currents, which is the quantity required in the measurement of extinction coefficients (see above).

It should be noted that, since the two compensation operations are simultaneous, the time  $t$  is necessarily exactly the same for both and vanishes from the final result. Thus the determinations do not involve the control or measurement of times of exposure. Also the method is essentially a complete null method. The potential of both the anode systems and of the electrometer needle is kept throughout practically at earth, so that leakage troubles are reduced to a minimum. Moreover, the voltage across both the photoelectric cells remains practically constant throughout and the greatest possible constancy of behaviour is thus ensured, although with vacuum cells at saturation the advantage of this is small.

In actual practice it is found impossible to cut off the light at the precise moment at which the slider of the potentiometer  $D$  has been brought to the end of its travel. Thus the electrometer needle usually shows a slight deflection after the light is cut off. To correct this small lack of compensation before moving over the switch  $S_1$ , use is made of the small trimming rheostat  $T$ , which is provided with lead-screw motion to allow of rapid operation. A small touch on this rheostat suffices to correct the compensation before changing

over  $S_1$  to the other system to see how nearly the dual compensation has been effected

According to what has been said so far, the determination of a photoelectric current ratio would involve a lengthy series of trials with different resistance ratios until the correct value was arrived at by a process of "bracketing" Actually, however, a method is adopted whereby the correct resistance ratio is determined directly An approximately correct value is first found by trial—this can be done quite rapidly—and then the electrometer deflection on changing over switch  $S_1$  is observed From these data the correct resistance ratio can at once be calculated

Let us denote by the subscript 1 the quantities associated with the photoelectric cell  $P_1$  (either  $P_s$  or  $P_m$ ) which gives the greater current under the conditions considered Quantities associated with the other cell  $P_2$  we designate by the subscript 2 At the correct resistance ratio for dual compensation, which we will take as  $5000/X$ , the quantities of charge  $Q_1$  and  $Q_2$  given respectively by the two cells  $P_1$  and  $P_2$  during the time of compensation would be  $-Q_1 = C_1 V$  and  $-Q_2 = C_2 V X / 5000$ , where  $V$  is the total potentiometer voltage applied Now consider an actual measurement in which a slightly incorrect value  $(X + \Delta X)$  is used instead of  $X$  Let the experimental procedure always be to compensate the system 2 by manipulation of the potentiometer slider and then, after cutting off the light, to switch over to system 1 (This is a necessary stipulation, as the procedure affects the considerations—see below) At the incorrect setting  $(X + \Delta X)$  we have

$$-Q'_2 = C_2 V \frac{X + \Delta X}{5000} = -Q_2 \left(1 + \frac{\Delta X}{X}\right)$$

Since the time of illumination of both photoelectric cells is necessarily the same, it follows that

$$-Q'_1 = -Q_1 \left(1 + \frac{\Delta X}{X}\right) = C_1 V \left(1 + \frac{\Delta X}{X}\right) \quad (1)$$

Thus the compensation of system 1 is imperfect, and on switching over the electrometer needle to this system from the perfectly compensated system 2 (after cutting off the light) an electrometer deflection will be observed Tests showed that the electrometer used gave a linear voltage response over the range of measurement, so that the observed deflection of  $n$  scale divisions corresponds to a needle potential of  $n/k$ , where  $k$  is the voltage sensitivity (We regard  $k$  as a positive constant, i.e., we make the sign of  $n$  the same as that of the needle potential.) The magnitude of the charge resident in the

induction condenser is now  $C_1 [V - (n/k)]$ , while that in the condenser formed by the needle system and the rest of the apparatus, is  $Z_1 (n/k)$ , where  $Z_1$  is the corresponding capacity. The net charge on the whole needle system is the photoelectric charge  $Q'_1$ , since the photoelectric cell is the only source. Hence taking account of signs, we have

$$-Q'_1 = C_1 \left( V - \frac{n}{k} \right) - Z_1 \frac{n}{k},$$

which together with (i) gives

$$\Delta X = - \frac{X}{C_1 V k} (C_1 + Z_1) n \quad (ii)$$

From this we see that the absolute error in  $\Delta X$  for a given error in the reading of  $n$  is directly proportional to  $X$ , i.e., the relative error is the same for all values of  $X$ . This advantage is not obtained when the procedure is to compensate the cell  $P_1$  giving the greater current and then to switch over to the cell  $P_2$ , for then a similar calculation shows that a given error in the reading of  $n$  corresponds to the same absolute error in  $\Delta X$  for all values  $X$ , i.e., the relative error increases as  $X$  decreases.

The correct resistance  $X$  is calculated from the value  $R$  actually used by means of the equation

$$X = \frac{R}{1 - \frac{(C_1 + Z_1)n}{C_1 V k}}, \quad (iii)$$

which is obtained from (ii) by substituting  $\Delta X = R - X$ . The value of  $V$  is read off from the voltmeter  $M$  (see figure) with the slider  $D$  at the end of its travel. Of course the slight readjustments of the trimming rheostat  $T$  cause corresponding changes in  $V$ . However, on account of the smallness of the resistance of  $T$  relative to that of  $D$ , the variations of  $V$  are only of the order of 1%, and since they produce errors in  $X$  of the second order of smallness, they are quite negligible. Thus  $V$  may be taken as constant throughout a run. The constant  $k$  of the electrometer is determined directly. Finally, considerations similar to those given above show that the remaining quantity  $(C_1 + Z_1)/C_1$ , which is a constant of the apparatus, can be determined once and for all as the ratio of the electrometer deflections in the following independent cases: (i) a certain voltage is applied to the needle direct, (ii) the needle being insulated, at zero potential and connected to system 1, the same voltage is applied to the induction condenser of capacity  $C_1$ . Thus the value of the constant  $K = (C_1 + Z_1)/(C_1 V k)$  can be determined and the correct values of  $X$  required in the extinction coefficient determinations can be calculated from

equation (iii) in the form  $X = R/(1 - Kn)$ . In actual observations the deflection  $n$  is always small and the quantity  $Kn$  is always small relative to unity. Thus the method is a true null method for the cell  $P_1$  and approximately a null method for the cell  $P_2$ .

### III Test of Apparatus and Method

In order to obtain an idea of the performance of the apparatus, three test runs were carried out with an N/15 aqueous solution of potassium nitrate. In each run determinations were made at seven different wave-lengths in the ultra-violet. The runs were performed on different days and with different samples of solution, so that the accuracy obtained is the over-all accuracy of the whole procedure, including beside the actual measurement the cleaning and filling of the absorption vessels, etc. It is known\* that Beer's law is obeyed up to concentrations greater than that used. The molecular extinction coefficients  $\epsilon$  given in Table I are calculated from the formula

$$\epsilon = \frac{1}{cd} \log_{10} \frac{I_0}{I},$$

where  $c$  is the concentration in mols per litre,  $d$  the thickness of the solution in centimetres, and  $I_0$  and  $I$  the incident and transmitted intensities respectively. Here  $c = 1/15,000$  and  $d = 2.000$ .

Table I—Determinations of Molecular Extinction Coefficients of  $KNO_3$

$\lambda$ in $m\mu$	313	302	297	289	280	265	254
$\epsilon$ (run I)	5.38	7.19	6.85	5.72	3.75	1.60	1.63
$\epsilon$ (run II)	5.38	7.21	6.88	5.67	3.77	1.56	3.56
$\epsilon$ (run III)	5.39	7.17	6.92	5.66	3.82	1.62	3.57
$\epsilon$ (mean)	5.38	7.19	6.88	5.68	3.78	1.59	3.59
% error (run I)	0	0	-0.4	+0.7	-0.8	+0.6	+1.1
% error (run II)	0	+0.3	0	-0.2	-0.3	-1.9	-0.8
% error (run III)	+0.2	-0.3	+0.4	-0.35	+1.1	1.9	-0.6
$\epsilon$ (von H. and R.)	5.26	6.93	6.84	5.61	3.68	1.59	3.59

The first row of Table I gives the wave-length in  $m\mu$  of the light used. The next three rows contain the results of the three independent runs, while in the fifth row is given the means of the values obtained. The next three rows

\* von Halban and Eisenbrand, 'Z. phys., Chem.', vol. 132, p. 401 (1928)

contain the percentage differences from the mean value. Since the errors of the method are due to small random changes in the apparatus, these figures give an idea not only of the reproducibility but also of the accuracy of the determinations. Thus in the table they are called errors.

The bottom row of the table contains the values of  $\epsilon$  taken from the data of von Halban and Eisenbrand\* upon the nitrate ion in dilute solution, which are reproduced here for the sake of comparison. The agreement with the mean values given in the fifth row of Table I is fair, but these mean values are definitely somewhat higher than the corresponding values given by von Halban and Eisenbrand.

It is seen from Table I that the maximum percentage deviation from the mean in the present determinations is almost always under  $\frac{1}{2}\%$  for the higher values of  $\epsilon$ . At the lower values of  $\epsilon$  the maximum error becomes greater, but is nowhere more than 2%. Since  $\epsilon$  is proportional to  $\log(I_0/I)$ , the percentage error in  $\epsilon$  corresponding to a given percentage error in the experimental determination of  $I_0/I$  is greater at small values of  $I_0/I$  than at larger values. This accounts for the greater percentage errors at the last three wave-lengths in Table I, and particularly at  $\lambda = 265 \text{ m}\mu$ . The corresponding errors in the measurements of  $I_0/I$  are actually about the same throughout (nowhere greater than 1.3%). In the determination of extinction coefficients it is clearly of advantage for the sake of accuracy to choose (when possible) the concentration and thickness of liquid so as to avoid values of  $I_0/I$  near to unity. With the apparatus here described, the limits of error in the determinations are then about  $\pm \frac{1}{2}\%$ .

This performance could probably be improved by the introduction of suitable modifications. Thus, referring to equation (iii), we see that the sensitivity can be increased (a) by increasing the compensating potential  $V$ , and (b) by using an induction condenser of greater capacity  $C_1$ , thereby decreasing the value of the quantity  $(C_1 + Z_1)/C_1$ . In (a) the possible increase of sensitivity is in principle unlimited, in (b)  $(C_1 + Z_1)/C_1$  approaches the limiting value unity as  $C_1$  approaches infinity. It should be noted, however, that in both (a) and (b) the increase of sensitivity is only obtained at the cost of lengthening the time required in compensation. Moreover, at higher sensitivities the value of  $R$  used must approximate more closely to the correct value  $X$ , otherwise the method is no longer approximately a null method. In the actual determinations quoted above the time of compensation was usually about  $\frac{1}{2}$  minute,

\* *Loc. cit.*, Table I, p. 406.

and the accuracy obtained was considered satisfactory for the particular purpose for which the arrangement was set up

The apparatus and method described above are to be used for investigating the ultra-violet absorption of certain extracts from blood sera. The writer wishes to express his thanks to the Nottingham Branch of the Imperial Cancer Campaign for a grant covering the cost of the apparatus

### *Summary*

A new type of photoelectric photometer is described, suitable for the measurement of extinction coefficients of substances in solution, more particularly in the ultra-violet region. Errors due to fluctuations of the light source are eliminated by the use of two photoelectric cells, while the simultaneous electrostatic compensation of both cells by means of one and the same potentiometer system does away with the necessity of measuring or controlling times of exposure. No devices for the quantitative variation of light intensity are required. The quantity measured is the ratio of the compensating potentials, and the actual observations are of a resistance and an electrometer throw. The method is tested by a series of determinations of the molecular extinction coefficient of potassium nitrate in dilute solution. The results show that the error in the measurement of  $I_0/I$  does not exceed 1.3%, which corresponds, under the conditions of experiment, to a maximum error of about 1% in the determination of extinction coefficients.

[*Note added in proof, January 2, 1934*]—Subsequent work with the instrument has shown that the approximation (iii) above, though otherwise sufficiently accurate, becomes inadequate when  $R$  is small relative to 5000. It is then no longer permissible to neglect the small induction effect of system 1 upon system 2. The equations  $-Q_1 = C_1V$  and  $-Q_2 = C_2VX/5000$  remain unaltered, but if  $c$  is the small capacity between the two systems during compensation, we have

$$-Q'_2 = \frac{C_2VR}{5000} + cv = -Q_2 \left( \frac{R}{X} + \frac{5000cv}{C_2VX} \right), \quad (iv)$$

where  $v$  is the small potential of system 1 at the end of the compensation operation and before switching over. Hence instead of (i) we obtain

$$-Q'_1 = -Q_1 \left( \frac{R}{X} + \frac{5000cv}{C_2VX} \right) = C_1V \left( \frac{R}{X} + \frac{5000cv}{C_2VX} \right) \quad (v)$$

At this stage

$$-Q'_1 - C_1(V - v) - cv - y_1v, \quad (vi)$$

where  $y_1$  is the capacity between system 1 and its earthed surroundings. In changing over switch  $S_1$  a charge  $-c_s v$  is transferred from system 2 to system 1,  $c_s$  being the small capacity between system 1 and the switch arm. As a result of the switch-over system 1 assumes a potential  $n/k$  (the corresponding electrometer throw  $n$  being observed), while system 2 assumes a potential  $p$  by induction. We now have

$$-(Q'_1 - c_s v) - C_1\left(V - \frac{n}{k}\right) + c\left(p - \frac{n}{k}\right) - z_1 \frac{n}{k}, \quad (vii)$$

where  $z_1$  is the new value corresponding to  $y_1$ . Also for system 2,

$$(C_2 + c + z_2)p = c\left(\frac{n}{k} - v\right), \quad (viii)$$

$c$  being the same in both positions of the switch.

Elimination of  $Q_1$ ,  $Q_2$ ,  $Q'_1$ ,  $Q'_2$ ,  $v$  and  $p$  from these equations gives an expression for the required quantity  $X$  in terms of constants of the apparatus and the observed quantities  $R$  and  $n$ . The result becomes identical with (iii) if we put  $c = c_s = 0$ , while for  $n = 0$  it reduces to  $X = R$ . Without reproducing the expression we may note that for any particular value of  $X$  it gives (as did the simpler treatment) a linear relationship between  $n$  and  $R$ . This is confirmed by experiment, and  $X$  may be conveniently determined by observing the electrometer throws for two different values of  $R$  and interpolating linearly to zero throw.

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# Changes in the Raman Spectrum of Sulphuric Acid on Dilution

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## Introduction

The results of previous workers upon the Raman effect of sulphuric acid show a considerable lack of agreement, due no doubt in part to the intensity of the accompanying continuous background and the apparent diffuseness of certain of the lines. Thus both Nis† and Woodward‡ found that change of the concentration of the acid greatly modified the appearance of the spectrum, influencing both the positions and the relative intensities of the lines. Woodward gave photometer curves of the spectra for concentrations ranging from 100% to 25% by volume, and attributed the observed intensity changes to the successive stages of ionization of the acid. The lines  $\Delta\nu = 910$  and  $1140\text{ cm}^{-1}$  (approximate values), which were strong in the 100% acid, became weaker and shifted somewhat as the concentration decreased, while a new line  $\Delta\nu = 1046\text{ cm}^{-1}$  made its appearance and increased correspondingly in intensity. At low concentrations this last line became double, a companion appearing at  $982\text{ cm}^{-1}$ . The frequencies 910 and  $1140\text{ cm}^{-1}$  were regarded as characteristic of the  $\text{H}_2\text{SO}_4$  molecule, 1046 of the  $\text{HSO}_4^-$  ion, and 982 of the  $\text{SO}_4^{--}$  ion. Other diffuse lines also showed frequency shifts. Bell and Fredrickson,§ however, whose measurements were made not only with a prism spectrograph but also with a grating, failed to find any appreciable shifts of the lines with dilution. Moreover, they stated that on their plates the 1046 line gave no indication of becoming double at low concentrations, but merely grew more diffuse. The findings of other workers are referred to in the discussion of our results (see below).

In view of the conflicting nature of the above evidence regarding the two stages of ionization of the acid, it was thought desirable to repeat the investigations using a higher dispersion than that employed by Woodward (*loc cit.*),

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† 'Jap. J. of Phys.,' vol. 5, p. 119 (1929), as quoted by Kohlrausch on p. 130 of "Der Smekal-Raman Effekt," published by Springer, Berlin, 1931

‡ 'Phys. Z.,' vol. 32, p. 212 (1931)

§ 'Phys. Rev.,' vol. 37, p. 1562 (1931)



the object of the work being to obtain more conclusive photometer curves of the spectra. The curves we have actually obtained not only give a clear picture of the intensity changes involved, but have also been used in a rational method of wave-length determination, the adoption of which has made it possible for us to propose a somewhat more detailed interpretation of the spectra than has hitherto been attempted.

### *Apparatus*

A modified form of Wood's well-known arrangement was used. In order to obtain comparable spectra it was essential that the alignment of the Raman tube and spectrograph should remain undisturbed throughout the series of exposures. The apparatus was therefore designed so as to permit of changing the liquid without in any way altering the position of the tube.

The main cylindrical part of the tube was surrounded with a thin water-jacket for cooling, and sloped at a small angle to the horizontal. At the upper end and immediately in front of the spectrograph slit was a plane window, and at the highest point of the tube a small air vent which could be tightly closed. The lower end of the tube was of the usual tapering horn-shape. The horn pointed downwards so that its end was the lowest point. The tube was filled and emptied from this end, which was attached to the middle of a relatively narrow vertical tube fitted with externally ground stoppers top and bottom. This observation vessel was made entirely of glass, the window being fused on. No grease was used on the ground stoppers.

The source of illumination was a 15-inch Cooper-Hewitt glass mercury lamp equipped with an automatic starting device whereby it could be switched on without being touched. This lamp was mounted close and parallel to the Raman tube, so that lamp and tube lay along the focal axes of a cylindrical chromium-plated mirror of elliptical cross-section closely surrounding them both. The water cooling maintained the liquid under investigation at a temperature between 20° and 25° C. during the exposures.

The spectrograph was a Hilger constant-deviation instrument with a dispersion of about 40 Å. per millimetre in the region investigated. (Compare 150 Å. per millimetre in the same region given by the spectrograph used by Woodward (*loc cit*) in obtaining his photometer curves.) Every precaution was taken to avoid stray light, screens and diaphragms being interposed at suitable points. In addition the whole instrument was enclosed in a blackened box from which only the collimator projected. The end of the collimator

carrying the slit was further surrounded by another blackened compartment totally enclosing the slit and the window of the Raman tube.

Both spectrograph and Raman tube were firmly mounted upon a single heavy base, the whole assembly being set up so as to slope at the small angle to the horizontal necessary for the proper functioning of the mercury lamp. Screw adjustments were provided for the important process of aligning the tube coaxially with the collimator. This was accomplished conveniently by sighting the inside of the tube from the focal plane of the spectrograph camera (*Schlussellochperspektive*). After careful alignment everything was firmly clamped in position and not disturbed in any way throughout the whole series of experiments.

### *Materials*

The solutions were made up from the pure acid (B D H "Pure 100%") and dust-free distilled water from a special still. Precautions were taken to exclude dust and, for the 100% acid, moisture. Following previous example, the concentrations used were 100%, 90%, 75%, 50%, 25%, and 10% by volume.

The photographic plates used (backed "Double-X-Press") were prepared by Messrs Ilford, Ltd., from an experimental emulsion. They combined a high gamma with a low inertia for the spectral region under consideration (4400-4600 Å.), and proved markedly superior to any other type of plate tested. They were developed in complete darkness for a uniform time of 4½ minutes, using a pyro-soda developer.

### *Experimental Method*

Preliminary exposures were taken with the 75% acid (which previous experiments had shown suffered most from background) in order to find the most suitable slit width and exposure time. Photometric measurements of the plates obtained showed that the ratio of the intensity of the Raman lines to that of the continuous background was practically constant for different slit widths over the range investigated (5 to 33  $\mu$ ). This result provides proof that the "lines" cannot be strictly monochromatic (see also below). As wide a slit as was compatible with the desired detail was therefore chosen (actual width 25  $\mu$ ). As is to be expected, it was found that for a given slit width the height of the lines above the background on the photometer curve increased with decrease of exposure time, provided that the background was not under-exposed.

In spite of the comparatively small numerical aperture of the spectrograph camera (1.9), the exposure times required were quite short. In order to obtain spectra of approximately comparable density, the exposure times for the different solutions were made inversely proportional to the volume concentrations. The most suitable exposure time for the 75% acid was first found by trial to be 10 minutes, and the others were then adjusted accordingly (viz., 7½, 8½, 10, 15, 30 and 75 minutes for the 100%, 90%, 75%, 50%, 25% and 10% solutions respectively). A shutter between the prism and camera objective was used for making the exposures, before each of which the mercury lamp was allowed to run for at least half an hour so as to attain a steady state.

The plates obtained were passed through a projection-type photoelectric microphotometer, electrometer readings being taken every 0.01 mm. of plate. The resulting photometer curves, which were plotted on a very large scale, are shown (much reduced) in fig. 1. Each plate was traversed from the 4339 to the 4916 Å mercury lines, but for the sake of clarity the full traverse including the mercury lines is reproduced only in the case of the 100% and 10% acids. In no case was the broadening of the exciting line 4358 Å sufficient to affect seriously the neighbouring 4339 Å line, which was always well separated.

It was concluded that the position of a line, as defined by its point of maximum intensity, could be determined with considerably greater accuracy from its shape in the photometer curve than by the adjustment of the cross-wires of a travelling microscope. Moreover, in using the photometer curves, due allowance could be made for the effect of the slope of the continuous background and the proximity of other diffuse lines upon the position of the line under consideration. Each spectrum was already provided with two satisfactory fiducial marks—the blue mercury line 4339 Å and the dark green mercury line 4916 Å. These were not over-exposed on any plate. They always appeared on the photometer curves as narrow lines with almost vertical sides, and their width corresponded exactly to the theoretical value for a monochromatic line, as calculated from the known widths of spectrograph and photometer slits (see below). No actual measurements upon the plates were necessary. The photometer curves were carefully plotted on millimetre graph paper using a very large scale and, when the true point of maximum intensity of a line had been decided upon, its position was read off directly. A dispersion curve on the same scale was constructed from the photometer curve of a mixed copper arc and mercury spectrum, photographed by means of a very small reflecting prism introduced between the spectrograph slit and the window of the Raman tube. The smooth dispersion curve passed exactly through every

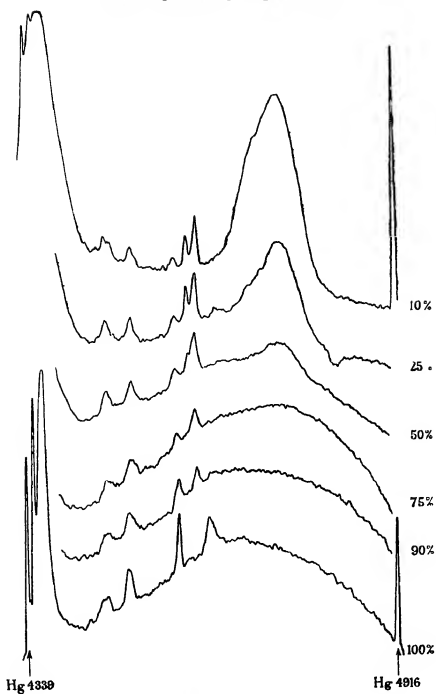


FIG 1

reliable known point, and the accuracy obtained with it was greater than that with which the positions of the lines could be determined. All the plates, including the comparison spectrum, were taken without disturbing the setting of the apparatus. In actual fact, although the plate-holder of the camera was necessarily moved each time, the distance on the plates between the two fiducial mercury lines was constant to within 1 part in 2000 (actual measurements 11 160, 11 155, 11 157, 11 160, 11 155, 11 160, and 11 160 mm on the 100%, 90%, , 10% acid and comparison plates respectively). Hence the readings taken from the different photometer curves could be immediately transferred to the single dispersion curve and the wave-lengths obtained directly. For conversion to wave-numbers use was made of Kayser's "Tabelle der Schwingungszahlen". The estimated error in the determination of the wave-length of a sharp line is less than  $\pm 0.2 \text{ \AA}$  (corresponding to  $\pm 1 \text{ cm}^{-1}$ ). Owing to the diffuseness of the Raman lines and the presence of the background, the  $\Delta\nu$ -values given below vary in accuracy between this and  $\pm 5 \text{ cm}^{-1}$  or more, and the probable error is given for each line individually.

### *Experimental Results and Discussion*

For convenience of comparison, the photometer curves of figs 1 and 2 have been arranged above one another so that their wave-length scales coincide to within  $0.2 \text{ \AA}$ . Fig 1 is a small-scale reproduction showing the general shape of the curves. It will be seen that the continuous background is in the form of a band extending from about 4400 to about 4900  $\text{\AA}$ . In studying the curves this background must not be confused with the well-known Raman band of water, which is much narrower and stands out most markedly in the most dilute acids (see below). The behaviour of the background is interesting. Its intensity at first increases as the concentration of the acid is reduced from 100%,\* but then drops off again in a striking manner. At 10% acid the background has become negligible, although the exposure time is ten times longer than for the 100% acid. This indicates clearly that the background cannot be wholly due to Rayleigh scattering of the continuous spectrum of the mercury lamp in this region, for the intensity of the classically scattered mercury lines is observed to increase regularly with the exposure time. Our curves would thus seem to lend support to the view of Placzek and van Wijk† that the background consists mainly of a true continuous Raman effect due to vibrations of molecular complexes.

\* It must be remembered that the exposure time increases in proportion to the dilution

† 'Z. Physik,' vol. 70, p. 287 (1931)

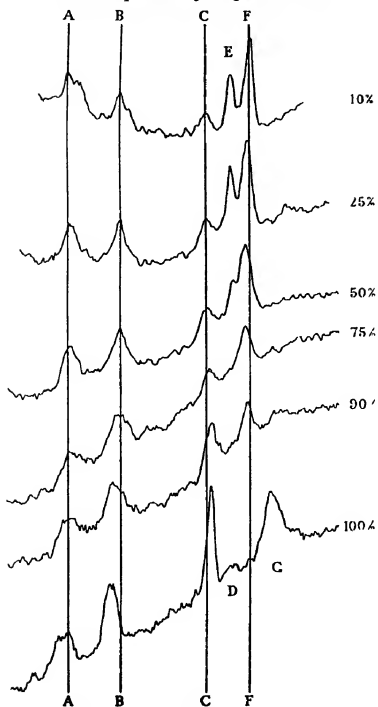


FIG 2.

The characteristically unsymmetrical Raman band of water, excited by 4047 Å, begins to show at 75% acid and increases rapidly in intensity as we pass to greater dilutions. At no stage is there any indication of its splitting into two separate components in the manner observed in the case of nitric acid (see Woodward, *loc cit*).

Fig. 2, on a rather larger scale, shows the Raman lines (all excited by 4358 Å) in more detail. The vertical lines drawn right down the figure pass through points of the same wave-length on the different curves. For convenience of discussion the Raman lines have been lettered A to G. It will be seen at once that the curves provide clear evidence in support of the general conclusions of Woodward (*loc cit*). The line F, supposed characteristic of the  $\text{HSO}_4^-$  ion, is completely absent in the 100% acid, but appears strongly in the 90% and increases in intensity with dilution. Already at 50% its companion E, ascribed to the  $\text{SO}_4^{2-}$  ion, has made its appearance, while in the lower concentrations it is stronger and quite clearly separated from the  $\text{HSO}_4^-$  line. The strong line G and the weak line D of the pure acid (the latter not very marked on the photometer curve, but visible on the plate) vanish rapidly on dilution; they are ascribed to the  $\text{H}_2\text{SO}_4$  molecule. The line D has not been found by previous workers.

In addition to the intensity changes outlined above, an apparent general wave-length shift of the lines with dilution is to be observed. The lines A, B, and F appear to move to longer wave-lengths as the acid concentration is reduced, while C shows the reverse behaviour (compare Nisi and Woodward, *loc cit*). Such shifts have also been noted by Specchia,\* who states, however, that all the lines are displaced to longer wave-lengths. On the contrary, Bell and Fredrickson (*loc cit*), using a grating, failed to find any appreciable changes of wave-length with dilution, except for a possible extremely small shift in the single case of  $\lambda$  4470 (B).

Our results would seem to leave no doubt as to the reality of the displacements under consideration. It appears difficult, however, to explain shifts of the observed magnitudes as due to alterations of the vibrational frequencies by intermolecular forces dependent upon the concentration. In this connection we may refer to some measurements of Embirkos† upon the frequency of the strongest line of the sulphate ion in salt solutions. For lithium and magnesium sulphates he found that this frequency increased with increase of concentration. For the latter salt he gave the values  $985.7 \text{ cm}^{-1}$  for 1 N solution and  $995.7$

\* 'Atti Accad. Lincei,' vol. 13, p. 754 (1931).

† 'Z. Physik,' vol. 65, p. 266 (1930).

$\text{cm}^{-1}$  for 2 N solution. We have carefully repeated the measurements upon this salt over an extended concentration range, but find no evidence whatever for a frequency shift of the magnitude reported by Embirikos. Our results are given in Table I, from which it is seen that the frequency of the line is independent of the concentration within the limits of error of our measurements.

Table I - Frequency of Strongest  $\text{SO}_4^{--}$  Line in Solutions of Magnesium Sulphate

Concentration of $\text{MgSO}_4$	$\Delta\nu$ (obs) in $\text{cm}^{-1}$
1.25 N	$982 \pm 2$
2.55 N	$982 \pm 2$
5.1 N	$981 \pm 2$

Returning now to the photometer curves for sulphuric acid given in fig. 2, it is seen on closer inspection that the wave-length shifts of the lines B and C are not gradual and uniform, but occur entirely in the region of 75% acid. The  $\Delta\nu$ -values, as determined from the points of maximum intensity of the lines, are given in Table II. Both lines are relatively sharp and of practically constant frequency both at the high and the low concentrations, but broaden and become less intense in the intermediate stages. The broadening occurs on the

Table II -- Frequencies of Points of Maximum Intensity of B and C

Acid concentration	Line B (in $\text{cm}^{-1}$ )	Line C (in $\text{cm}^{-1}$ )
%		
10	$\left\{ \begin{array}{l} 596.5 \\ 594.5 \\ 594.5 \\ 590 \\ 583 \\ 555.5 \end{array} \right.$	$\left\{ \begin{array}{l} 896 \\ 894.5 \\ 896 \\ 903 \\ 909.5 \\ 910 \end{array} \right.$
25		
50		
75		
90		
100		

side towards which the frequency is apparently changing, thus giving rise to the intermediate values given in Table II. These facts would suggest that in each of B and C we are dealing with two independent frequencies of practically constant value. In each the one which is predominant at 100% rapidly diminishes in intensity with dilution, while the other becomes correspondingly more intense, the former would naturally be regarded as characteristic of the  $\text{H}_2\text{SO}_4$  molecule, the latter of the  $\text{HSO}_4^-$  ion. Actually it was found that for all the apparent



intermediate frequencies recorded in Table II the shape of the broadened line in the photometer curves is indeed consistent with an analysis into two relatively narrow, symmetrically shaped components having the practically constant frequencies found in the most concentrated and most dilute acids respectively. The idea here proposed is illustrated diagrammatically in fig 3, which is a tentative line scheme of the spectra. The vertical lines in the diagram represent relatively narrow components or lines of practically constant frequency (except for the line F—see below), and the course of the intensity changes is indicated for each by corresponding variations in height.

The "line" A is slightly more complicated. Comparison with the widths of B and C shows at once that A is relatively broad throughout the whole range of concentrations. It must either be a diffuse band or it must consist of more than one component frequency. In the light of what has been said above about the lines B and C, the nature of the changes of shape of A with dilution (see fig 3) suggests an analysis into three symmetrically shaped and relatively narrow components of practically constant frequency. These are represented diagrammatically in fig 3, which also shows qualitatively the intensity trend of each. Thus the behaviour of A is explicable on the view that it is made up of a central component of approximately constant intensity at all concentrations, a second component of lower frequency, which dies away from 100% to 75% acid, and a third of higher frequency which first appears at the intermediate concentrations and becomes stronger with dilution. This explanation finds support in the fact that on the photographic plate for the 100% acid the line A appears to the eye to be double (contrast effect) \*.

As to the line E it was found that, after correction for the influence of the neighbouring line F, a constant value of  $\Delta\nu$  was obtained within the estimated limits of error.

The line F, however, seems to be an exception to the rule of practically constant frequency. It shows a frequency shift greater than the estimated maximum error of determination (see Table III below, which gives the measured values for all the lines or proposed components). A real displacement of F towards higher frequency in the lowest concentrations seems established. It may be noted that this line has a peculiar unsymmetrical shape and an excessive breadth at intermediate concentrations. This may perhaps mean

\* This double appearance is also noted by M $\acute{e}$ lard in a paper ('C R Acad. Sci. Paris,' vol. 197, p. 582 (1933)) on the Raman effect of 100% sulphuric acid, published since the conclusion of our experiments and the writing of this account. His experimental findings are everywhere in harmony with our own.

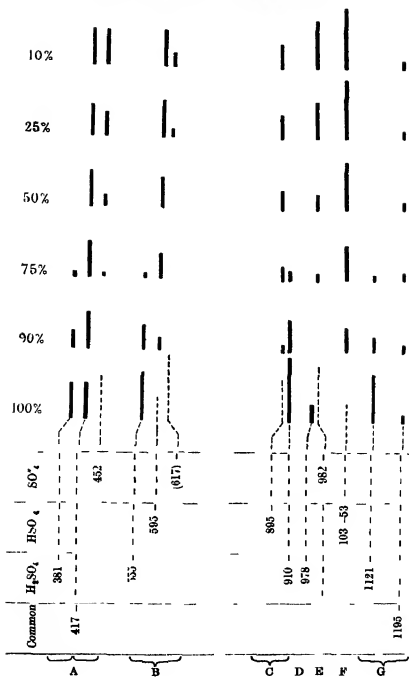


FIG. 3.

Table III—Observed Frequencies (in  $\text{cm}^{-1}$ )

	A			B			C			D	E	F	G	
10%	—	418 $\pm 5$	454 $\pm 8$	—	606.5 $\pm 2$	(617)	896 $\pm 5$	—	—	—	983 $\pm 2$	1053 $\pm 2$	—	—
25%	—	(418)	450 $\pm 6$	—	504.5 $\pm 2$	—	894.5 $\pm 3$	—	—	—	981 $\pm 3$	1043 $\pm 3$	—	—
50%	—	(414)	(451)	—	594.5 $\pm 3$	—	896 $\pm 5$	—	—	—	(985)	1036 $\pm 4$	—	—
75%	(381)	417 $\pm 5$	—	(555)	(595)	—	(894)	(910)	—	—	—	1035 $\pm 4$	—	—
90%	(381)	(414)	—	(555)	(595)	—	(896)	909.5 $\pm 3$	—	—	—	1038 $\pm 4$	—	—
100%	381 $\pm 6$	417 $\pm 5$	—	555.5 $\pm 2$	—	—	—	910 $\pm 2$	978 $\pm 4$	—	—	—	1121 $\pm 7$	(1165) $\pm 15$
Mean values	381	417	453	555.5	595	(617)	895.5	910	978	982	1036 to 1053	1121	(1186)	(1186)

that we are dealing with more than one component, but the evidence is too slight to justify further discussion

The strong line G ( $1121\text{ cm}^{-1}$ ) in the spectrum of the 100% acid rapidly disappears on dilution. In the photometer curve it is seen to be definitely broadened on the long-wave side, and we find that this broadening can be interpreted as due to the overlap of a weak diffuse line with its centre at about  $1195\text{ cm}^{-1}$ . Although this line does not show up on the photometer curves (except perhaps feebly in the 25% curve), it is actually visible by eye on all the plates except those for 90% and 75% acid which suffer most from background.

The  $\Delta\nu$ -values given in Table III are those of the lines or components referred to in the above discussion. Those components not unambiguously defined by some part of the smoothed photometer curves are placed in brackets, because of their uncertainty they have not been used in reckoning the mean values.

Usually the course of the intensity changes with dilution, fig. 3, indicates the molecular species ( $\text{H}_2\text{SO}_4$ ,  $\text{HSO}_4^-$ , or  $\text{SO}_4^{2-}$ ) of which any given line is characteristic. Thus the frequencies 381, 555, 910 (strong), 978 (weak) and  $1121\text{ cm}^{-1}$  are present in the pure acid, but die out by 75%. They are therefore presumably characteristic of the  $\text{H}_2\text{SO}_4$  molecule. (The frequency 978 here referred to must not be confused with the fortuitously almost equal frequency 982, which appears only in the most dilute solutions and is characteristic of the  $\text{SO}_4^{2-}$  ion.) The lines 595, 895, and  $1036\text{--}1053\text{ cm}^{-1}$  are not present at all in the pure acid, but appear and rapidly become stronger on dilution, reaching maximum intensity in the neighbourhood of 25% acid. This is the behaviour to be expected of lines characteristic of the  $\text{HSO}_4^-$  ion. Nisi (*loc cit*) described the last of these frequencies (the strongest) as present in the pure acid (compare also Fadda\*). Since even a small addition of water suffices to bring out this line (by 90% acid it is already strong), probably this discrepancy is to be explained as due to traces of water in the supposedly 100% acid of Nisi. Returning to our own results, the lines 452 and  $982\text{ cm}^{-1}$ , although present in the 50% acid, and possibly even in the 75%, do not become markedly appreciable until the acid content has been reduced to 25%, and are strongest in the lowest concentrations. This behaviour naturally supports the view that these lines are characteristic of the  $\text{SO}_4^{2-}$  ion. These allocations of the different frequencies are indicated beneath the line scheme of fig. 3.

The above views as to the origins of the different lines find confirmation in the Raman spectra of solutions of normal and acid sulphates. A large number

\* 'Nuovo Cimento,' vol. 9, p. 1 (1932)

of workers\* agree in giving the following frequencies as characteristic of the  $\text{SO}_4^{2-}$  ion 452, 617, 982 (strong), and 1106 Of these 982 appears as a strong separate line (E) and 452 as a component of A in the most dilute acids Inspection of the 10% photometer curve at the point corresponding to 617 (namely, just on the long-wave side of B) gives faint indications of the presence of this frequency also Then of the frequencies 595, 895, and 1036-1053 (strong), which we regard as characteristic of the  $\text{HSO}_4^-$  ion, the first and the last have been observed by Woodward (*loc cit*) in a solution of  $\text{KHSO}_4$ , where they were accompanied by the  $\text{SO}_4^{2-}$  ion frequency 982 owing to the partial ionization of the  $\text{HSO}_4^-$  ions.

There still remain two lines of our spectra as yet unassigned—a moderately strong one at  $417\text{ cm}^{-1}$  and a doubtful diffuse one of very low intensity at about 1195 Since the intensities of both these lines appear to remain approximately constant at all dilutions, they cannot be regarded as characteristic of any one of the species  $\text{H}_2\text{SO}_4$ ,  $\text{HSO}_4^-$ , or  $\text{SO}_4^{2-}$  in particular Their behaviour is, however, consistent with their being common to all three In this connection it is suggestive that approximately the same two frequencies occur in the Raman spectra† of other compounds whose formulæ may be written with two oxygen atoms linked to hexavalent sulphur, viz., sulphuryl chloride (408, 1182), chlorosulphonic acid (410, 1183), dimethyl sulphite (410, 1200) and diethyl sulphite (410, 1204) Moreover, these two frequencies seem to be the only ones common to all these molecules

In conclusion, we should like to make it clear that our analysis of the more diffuse Raman lines of sulphuric acid into relatively narrow components of practically constant frequency is put forward with some reserve While its correctness as an interpretation cannot be regarded as conclusively proved, it is nowhere in conflict with observed fact and the weight of supporting evidence makes it highly plausible It should be noted that our view does not exclude the possibility of relatively small frequency shifts (of the order of  $\pm 5\text{ cm}^{-1}$ ) with dilution Another point to be observed is that even the proposed components (like the simple Raman "lines") are only approximately monochromatic Each of them is broader than the theoretical value to be expected from the known widths of the spectrograph and photometer slits. This would be 0.18 mm of plate, and the mercury line 4916 Å, although more intense than any of the Raman lines, does indeed possess this breadth The Raman components, on the contrary, have an average breadth of about 0.30 mm. of

\* See, for example, the collected data in Kohlrausch, "Der Smekal Raman-Effekt"

† Matcoski and Aderhold, 'Z. Physik,' vol. 68, p. 683 (1931)

plate. The excess corresponds to an actual line breadth of about 5 Å or 25  $\text{cm}^{-1}$ . This result is in harmony with the conclusions from the preliminary experiments with various slit widths (see above), which give 8  $\text{cm}^{-1}$  (corresponding to the widest slit used) as a lower limit for the breadth of the Raman lines.

### Summary

Microphotometer curves of the Raman spectra of sulphuric acid are reproduced for concentrations ranging from 100% to 10%. The intensity changes of certain of the lines give a clear picture of the two stages of ionization of the acid. Thus on dilution the frequencies 910, 978, and 1121  $\text{cm}^{-1}$  (characteristic of the  $\text{H}_2\text{SO}_4$  molecule and present in the pure acid) rapidly vanish, while the frequency 1036 (characteristic of the  $\text{HSO}_4^-$  ion and absent in the pure acid) makes its appearance and increases in intensity. At 50% acid another frequency 982 (characteristic of the  $\text{SO}_4^{2-}$  ion) appears and subsequently becomes stronger. Other diffuse lines are found to change somewhat in frequency with dilution, but it is shown that their behaviour is consistent with an analysis into relatively narrow components of practically constant frequency, ascribable to the different molecular species present. On this view the characteristic frequencies come out as  $\text{H}_2\text{SO}_4$ , 381, 555, 5, 910, 978 and 1121,  $\text{HSO}_4^-$ , 595, 895, 5 and 1036,  $\text{SO}_4^{2-}$ , 452 and 982, while 417 and 1195  $\text{cm}^{-1}$  are common. The inherent breadth of the lines is found to be about 25  $\text{cm}^{-1}$ . Reference is also made to the behaviour of the continuous background. Measurements on solutions of magnesium sulphate give no evidence for a variation of the sulphate ion frequency 982 with concentration.

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*A Comparison of Experiment and Calculated Wave-Profiles and Wave-Resistances for a form having Parabolic Waterlines*

By W C S WIGLEY

(Communicated by T H Havelock, F R S—Received October 4 1933)

*Introduction*

It has been found possible to calculate the wave-resistance of simple three dimensional forms, and a number of such results have been obtained\* which on the whole have shown good agreement with actual measurements when the forms are such as not to violate the assumptions on which the calculations are founded. But the calculation of wave-profiles is at present limited to two dimensional forms having infinite draft. Hence for the purpose of this research two models were made, at the William Froude Laboratory, one of the maximum draft possible under mechanical limitations, with vertical sides and a flat bottom, the other having the greatest draft and displacement which would allow of the measurement of its resistance over the desired range of speeds, but with vertical sides for only half its draft and being then rounded off in vertical section to an edge at the keel. The first model (No 1254) was used to measure wave-profiles which were compared with those calculated for a model of infinite draft, the second model (No 1302) was used for the comparison of measured and calculated resistances.

It was impossible to use the first model for this purpose as the added resistance due to eddying round the sharp corners at the bilges would render quite meaningless any calculation of the frictional resistance which must be subtracted from the measured force to find the wave-resistance. Its wave-resistance was, however, calculated in order to show the similarity in shape between the calculated resistance curves of the two models. Fig 1 shows perspective sketches of these models and the equations of their surfaces appear below (equations (1) and (7) respectively)

*List of Symbols used throughout the Paper*

- $L$  = total length of model,
- $l$  = length of curved entrance or run,
- $2a$  = length of parallel body,

\* See 'Proc. 3rd Int. Cong. Appl. Math.', Stockholm, 1930, vol 1, p 58, for a summary of these results up to 1930

$2b$  = maximum beam,

$d$  = draft,

$c$  = speed of advance,

$\rho$  = density of water, taken as 1.94 foot lb sec units in calculations,

$g$  = acceleration due to gravity, taken as 32.2 in calculations,

$\kappa = g/c^2$ ,

$\zeta_w$  = surface elevation due to wave-system,

$\zeta_i$  = surface elevation due to "non-wave" portion of the disturbance caused by the motion of the form,

$R_w$  = resistance due to wave-making,

$H_0$  = Struve's function of order zero,

$Y_0$  = Bessel's function of the second kind and of order zero } (G. N. Watson's definitions)

$P_n$  and  $Q_n$  represent certain functions defined as they occur in the paper

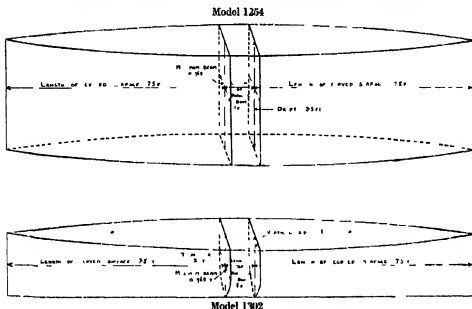


FIG. 1.—Sketch of under water forms of models 1254 and 1302

Foot-pound-second units and Cartesian axes were used throughout the calculations,  $Ox$  being in the direction of motion,  $Oy$  vertically downwards, and  $Oz$  at right angles to the other two axes. The origin of co-ordinates is amidships, in the centre line and in the plane of the undisturbed water surface, except that in equation (2) (to avoid negative values of the arguments of



certain functions) the origin has been taken at the bow and  $Ox$  reversed in direction

The calculations of resistances and wave-profiles are made on the following assumptions, by the methods either of Michell or of Havelock —

(1) The usual assumption in dealing with wave motion that the wave height is small compared with the other lengths occurring in the problem and that the velocities due to the wave motion are small compared with the other velocities involved. For ship waves this means that the wave height is small compared with the length and draft of the ship, and the velocities due to the wave motion are small compared with the ship's speed. It is also assumed that the angle made by the tangent plane to the side of the form with the vertical median plane of the ship is small.

(2) It is assumed that the effects of turbulence and viscosity can be neglected.

(3) In comparing calculated resistances with those of models in the tank, it is necessary also to assume that the alteration in the trim and sinkage of the hull during motion does not alter the effective wave-making form sufficiently to affect the wave motion appreciably. When wave-profiles alone have been compared this question does not arise, since the models were rigidly attached to the tank carriage and not allowed to trim or sink.

### *Measurement of Wave-Profiles*

The model (No 1254) used for the wave-profile measurements had water lines with the equation

$$\left\{ \begin{array}{ll} y = b \{1 - [\overline{x - a/l}]^2\}, & \text{from } x = a \text{ to } x = l + a \\ y = b, & \text{from } x = -a \text{ to } x = +a \\ y = b \{1 - [\overline{x + a/l}]^2\}, & \text{from } x = -a - l \text{ to } x = -a \end{array} \right\}, \quad (1)$$

where

$$l = 7.5,$$

$$b = 0.484,$$

$$a = 0.5$$

The draft of the actual model was 3.5 feet, with 0.5 feet of necessary free-board. This gave a total depth of 4.0 feet, which was the maximum vertical dimension of a model which could be lifted into the Yarrow tank \*

\* The dimensions of the cross-section of the Yarrow tank are 30 feet wide by  $12\frac{1}{2}$  feet deep.

For the purpose of the measurement of the wave profiles this model was rigidly attached to the travelling carriage of the Yarrow tank, and was towed through the water at a series of speeds from 3.5 to 10.8 feet per second.

A network of vertical and horizontal lines had been previously marked on one side of the model, the horizontal lines being  $\frac{1}{4}$  inch apart, and the verticals normally 1 foot apart, but placed more nearly together near the ends of the model. During experiments at each speed the height of the water surface at each vertical was observed, the closer spacing near the ends of the model enabling more accurate observation where the slope of the water surface was greatest.

Two wooden beams were clamped to the carriage, forming extensions of 20 feet aft and 6 feet forward respectively of the centre line of the model. Each of these beams carried, at intervals of 1 foot, pointers sliding vertically which were adjusted during each experiment so that their pointed ends just touched the water surface. By this means the wave-profile could be plotted for a distance corresponding to the length of the beams. Additional pointers were again fitted where the slope of the wave-profile was steepest, i.e., near the bow and stern of the model. Owing to a very slow, long and persistent wave which is always set up in the tank during experiments, errors of measurement of  $\pm 0.05$  inch are unavoidable. Once an experiment has been made at one of the higher speeds, changes of this order in the water level are known to last for some 24 hours, and it is therefore impracticable to wait for their cessation.

#### *Calculation of Wave-Profiles*

Calculations were made for a form as given by equations (1) but having infinite depth using a formula given by Havelock† for the wave-profile of any two-dimensional form of infinite draft. The profile actually given by the formula is that along the centre line of the form (the axis of  $x$ ), but the error involved in using this formula to find the profile on the curved surface of the model (which is of the same order as those introduced by the assumption that the angle is small between the tangent plane to the side of the form and the median plane) can be neglected. For this particular form the expressions for the "wave" and "non-wave" portions of the profile become respectively

$$\zeta_w = (8b/\pi\kappa l) [P_0\{\kappa x\} + P_0\{\kappa(x - 2l - 2a)\} - (1/\kappa l) (P_0^{-1}\{\kappa x\}) \\ - P_0^{-1}\{\kappa(x - l)\} + P_0^{-1}\{\kappa(x - l - 2a)\} - P_0^{-1}\{\kappa(x - 2l - 2a)\}], \quad (2A)$$

\* 'Proc. Roy. Soc., A, vol. 135, p. 11 (1932)

$$\zeta_1 = (-2b/\pi\kappa l) [Q_0\{\kappa x\} + Q_0\{\kappa(x-2l-2a)\} - (1/\kappa l)(Q_1\{\kappa x\}) \\ - Q_1\{\kappa(x-l)\} + Q_1\{\kappa(x-l-2a)\} - Q_1\{\kappa(x-2l-2a)\}] \quad (2B)$$

In these expressions the origin has been moved to the bow, and the positive direction of  $x$  is now astern. The functions  $P_0$  and  $P_1$  are supposed to be zero for negative values of their arguments, and for other values to be defined by the equations

$$\left. \begin{aligned} P_0(u) &= \int_0^{\pi/2} \sin(u \sec \phi) d\phi \\ P_0^{-1}(u) &= 1 + P_1(u) = 1 - \int_0^{\pi/2} \cos \phi \cos(u \sec \phi) d\phi \end{aligned} \right\} \quad (3A)^*$$

The functions  $Q_0$ ,  $Q_1$  are defined, for positive values of the arguments by the equations

$$\left. \begin{aligned} Q_0(u) &= \frac{\pi}{2} \int_0^u \{H_0(t) - Y_0(t)\} dt \\ Q_1(u) &= \int_0^u Q_0(t) dt \end{aligned} \right\}, \quad (3B)$$

where  $H_0$  is Struve's function and  $Y_0$  the Bessel function of the second kind, both of order zero, and by the convention that  $Q_0(-u) = -Q_0(u)$  and  $Q_1(-u) = -Q_1(u)$  for negative values of the argument.

Detailed calculations of the wave heights have been made for this form from equations (2) for seven speeds covering a range which corresponds with possible ship speeds. These speeds are given in Table I, with the value of the non-dimensional speed unit used in this paper, the (P) speed unit commonly used in shipbuilding work in this country and the so-called Froude number ( $c/\sqrt{gl}$ ) which is in common use abroad.

Table I

$c$	$c/\sqrt{gl}$	(P)	$c/\sqrt{gl}$ → (Froude number)
4.22	0.272	0.562	0.193
5.45	0.351	0.726	0.250
6.74	0.433	0.897	0.308
7.95	0.512	1.059	0.364
9.18	0.592	1.225	0.421
10.42	0.675	1.399	0.480
10.70	0.694	1.434	0.494

\* For further particulars, series, and asymptotic expansion of these functions, see Havelock, 'Proc Roy Soc,' A, vol. 108, pp 81, 82 (1925), vol 103, pp 577-579 (1923)



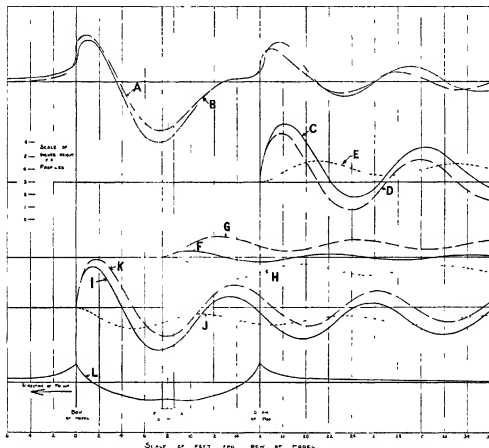


FIG. 3.—Model 1264. Analysis into components of calculated wave-profile. Speed 7.95 feet per second,  $\epsilon \sqrt{g l} = 0.512$

A. Observed wave profile

B. Total calculated wave-profile  $\zeta_w + \zeta_l$  given in (2a)

C. Total stern wave profile given by terms  $(8b/\pi\omega l) [P_0 \{ \epsilon(x-2l-2a) \} + (1/\omega) P_0^{-1} \{ \epsilon(x-2l-2a) \}]$  in (2a)

D. Profile of wave system commencing at stern due to stern angle given by term  $(8b/\pi\omega l^2) P_0^{-1} \{ \epsilon(x-2l-2a) \}$  in (2a)

E. Profile of wave system commencing at stern due to curvature of form given by term  $(8b/\pi\omega^2 l^2) P_0^{-1} \{ \epsilon(x-2l-2a) \}$  in (2a)

F. Sum of profiles of forward and after shoulder wave systems given by term  $(8b/\pi\omega^2 l^2) [P_0^{-1} \{ \epsilon(x-l) \} - P_0^{-1} \{ \epsilon(x-l-2a) \}]$  in (2a)

G. Profile of wave-system commencing at forward shoulder and due to curvature of form, given by term  $(8b/\pi\omega^2 l^2) P_0^{-1} \{ \epsilon(x-l) \}$  in (2a)

H. Profile of wave-system commencing at after shoulder and due to curvature of form, given by term

$$(-8b/\pi\omega^2 l^2) P_0^{-1} \{ \epsilon(x-l-2a) \} \text{ in (2a)}$$

I. Total bow wave profile given by terms  $(8b/\pi\omega l) [P_0 \{ \epsilon x \} - (1/\omega) P_0^{-1} \{ \epsilon x \}]$  in (2a)

J. Profile of wave-system commencing at bow and due to curvature of form given by term  $(-8b/\pi\omega^2 l^2) P_0^{-1} \{ \epsilon x \}$  in (2a)

K. Profile of wave-system commencing at bow due to bow angle given by term  $(8b/\pi\omega l^2) P_0 \{ \epsilon x \}$  in (2a)

L. Profile of symmetrical disturbance =  $\zeta_s$ , given in (2a)

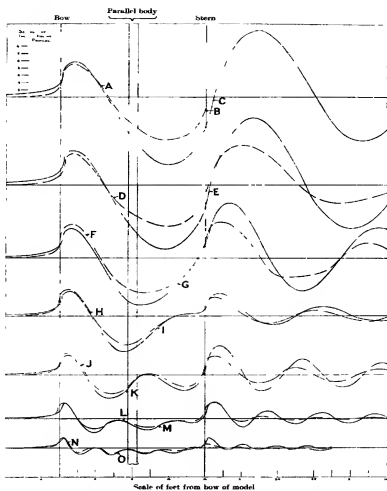


FIG. 2.—Model 1254. Comparison of calculated and observed wave profiles

- A Broken line is observed wave profile at 10.78 feet per second ( $c/\sqrt{g\lambda} = 0.694$ ) (No experimental values of wave height are available aft of the point marked B for this speed.)  
 C Full line is calculated wave profile at 10.78 feet per second ( $c/\sqrt{g\lambda} = 0.694$ )  
 D Broken line is observed wave profile at 10.50 feet per second ( $c/\sqrt{g\lambda} = 0.673$ )  
 E Full line is calculated wave profile at 10.50 feet per second ( $c/\sqrt{g\lambda} = 0.673$ )  
 F Broken line is observed wave profile at 9.20 feet per second ( $c/\sqrt{g\lambda} = 0.592$ )  
 G Full line is calculated wave profile at 9.20 feet per second ( $c/\sqrt{g\lambda} = 0.592$ )  
 H Broken line is observed wave profile at 7.95 feet per second ( $c/\sqrt{g\lambda} = 0.512$ )  
 I Full line is calculated wave profile at 7.95 feet per second ( $c/\sqrt{g\lambda} = 0.512$ )  
 J Broken line is observed wave profile at 6.74 feet per second ( $c/\sqrt{g\lambda} = 0.433$ )  
 K Full line is calculated wave profile at 6.74 feet per second ( $c/\sqrt{g\lambda} = 0.433$ )  
 L Broken line is observed wave profile at 5.45 feet per second ( $c/\sqrt{g\lambda} = 0.351$ )  
 M Full line is calculated wave profile at 5.45 feet per second ( $c/\sqrt{g\lambda} = 0.351$ )  
 N Broken line is observed wave profile at 4.22 feet per second ( $c/\sqrt{g\lambda} = 0.272$ )  
 O Full line is calculated wave profile at 4.22 feet per second ( $c/\sqrt{g\lambda} = 0.272$ )



The actual and the calculated wave-profiles for each speed are shown in fig 2. It will be seen that along the side of the form the chief peculiarities of the observed profiles are well reproduced by calculation. The observed bow wave is generally a little higher than the calculated, except at the highest speeds, whereas the stern wave is generally lower. Ahead of the form the disturbance is never so high as calculated, while aft of the form the disturbance (as would be expected) dies away much more quickly particularly at the slow speeds than it would in a perfect fluid.

Examination of the first equation (2) shows that the wave profile may be divided into six components, two originating at the bow, two at the stern, and one at each end of the parallel body, that is, at the forward and after shoulders. The two terms involving the function  $P_0$  are the same as would occur to produce bow and stern wave-systems in a form composed of straight lines having the same angle at bow or stern. This bow wave is decreased and this stern wave increased by the terms

$$-(1/\kappa l) P_0^{-1}(\kappa x) \quad \text{and} \quad (1/\kappa l) P_0^{-1}\{\kappa(x - 2l - 2a)\},$$

respectively, which may be considered as representing additional wave-systems due to the curvature of the form starting at bow and stern respectively. Owing to the presence of an extra factor  $\kappa$  in the denominator, these terms increase more rapidly with the speed than the terms in  $P_0$ , and their effect is therefore much greater at the higher speeds. The other two terms,

$$(1/\kappa l) P_0^{-1}\{\kappa(x - l)\} \quad \text{and} \quad (-1/\kappa l) P_0^{-1}\{\kappa(x - l - 2a)\}$$

represent wave-systems starting at the beginning and end of the parallel body, they also are of little importance at low speeds, owing to the factor  $\kappa$  in their denominator, and since they are equal in amplitude and opposite in phase and separated only by the length of parallel body—which corresponds, as the speed increases, to less and less difference in phase between them—their combined effect does not grow greatly with speed, and they are never of very great importance.

These statements are consistent with the known experimental facts, that the wave-making of a parabolic form is largely concentrated at the ends, and that it is an efficient form for low speeds, where the two terms mentioned in the second place above are of no great importance. In order to exhibit the relative importance of these terms, fig 3 has been drawn showing their respective sizes for  $c = 7.95$  ft per sec ( $c/\sqrt{gl} = 0.512$ ). To see the relative importance at any other speed it should be noted that the two systems depending on the bow and stern angles respectively will increase in height as the square of the



speed, while the remaining four systems will increase in height as the fourth power of the speed, and the wave-lengths of all the systems will increase in proportion to the square of the speed

From what has been written above it will be seen that the total bow wave-profile is given by the equation

$$\zeta_{Bow} = (4h/\pi\kappa l) [P_0\{\kappa x\} - (1/\kappa l) P_0^{-1}\{\kappa x\}], \quad (4)$$

and that the crests and troughs of the bow wave-system will therefore occur at positions given by values of  $x$  such that

$$\delta\zeta_{Bow}/\delta x = 0,$$

or, differentiating,

$$Y_0(\kappa x)/P_0(\kappa x) = -2c^2/\pi g l, \quad (5)$$

since  $d P_0(u)/du = -\pi Y_0(u)/2$  and  $d P_0^{-1}(u)/du = P_0(u)$ . Curves of the function  $Y_0(u)/P_0(u)$  have been plotted and with their aid the values of  $x$  satisfying equation (5) for a series of values of  $c$  have been calculated. A similar equation, with only the sign changed, holds for the stern wave system, curves showing the position of the crests and troughs of the waves of the two systems over a range of speeds have been calculated and plotted in fig. 4 on a base of  $c/\sqrt{gl}$ .

From these curves the speeds of coincidence in the crests and troughs of these systems can be seen, the relation of these to the maxima and minima of wave resistance will be discussed at a later stage in the paper.

#### *Calculation of Resistance for the Form of Great Draft*

The wave resistance of this form was calculated by the use of Havelock's result, which in the notation of this paper is given by the equation

$$R_w/c^2 = [64\rho b^2/\pi\kappa^2 l] \{ [1/3 + 1/\kappa^2 l^2] + \{P_2(p_1)/2 - P_4(p_1)/p + P_6(p_1)/2p^2\} - \{P_4(p)/p - P_6(p)/p^2 - P_4(p_2)/p + P_6(p_2)/p^2 - P_6(p_2)/2p^2\} \}, \quad (6)$$

in which, for brevity,  $p$  has been written for  $\kappa l$ ,  $p_1$  for  $2(l+a)\kappa$  or  $l\kappa$ ,  $p_2$  for  $(2a+l)\kappa$  and  $p_3$  for  $2a\kappa$ , whilst

$$\left. \begin{aligned} P_2(u) &= \int_0^{u/2} \cos^2 \phi \cos(u \sec \phi) d\phi \\ P_4(u) &= \int_0^{u/2} \cos^4 \phi \sin(u \sec \phi) d\phi \\ P_6(u) &= - \int_0^{u/2} \cos^5 \phi \cos(u \sec \phi) d\phi \end{aligned} \right\} *$$

\* See footnote \* on p. 148.



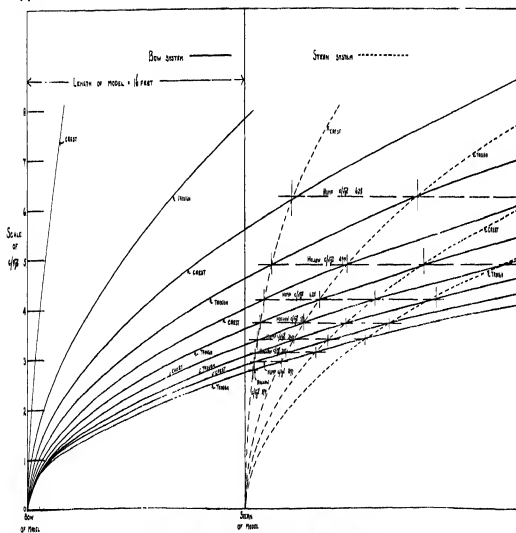


FIG. 4.—Model 1934. Positions of resist and lengths of bow and stern wave systems at various speeds.



Of the three terms of equation (6) in twisted brackets the first clearly represents the steady increase of wave making resistance with speed, apart from any question of interference between the wave-systems. The second, depending directly on the total length of the form, represents the interference between the bow and stern systems. Each of the terms in the third bracket represents some form of interference between the systems at the ends of the parallel body and the bow and stern systems, or between the systems at the ends of the parallel body themselves. These terms have not been considered separately because in this form their total effect is comparatively small.

Fig. 5 gives curves showing these three components, and also a curve of the total value of  $R_w/c^2$ . Owing to the great increase of this quantity with speed, the scale of the lower portions of the curves had to be exaggerated as shown in the diagram. It is clear that the main peculiarities of the total curve of  $R_w/c^2$  are due to the component corresponding with the second term in twisted brackets on the right-hand side of equation (6), and are therefore due to interference between the waves of the bow and stern systems. It is therefore of interest to compare the speeds of coincidence of crests and troughs of

Table II.—Values of  $c/\sqrt{gl}$  Corresponding to certain Phenomena

Model 1254			Model 1302		Maximum or minimum of $R_w/c^2$
Calculated values of $c/\sqrt{gl}$ for wave coincidences between bow and stern systems	Calculated values of $c/\sqrt{gl}$ for maxima and minima of		Values of $c/\sqrt{gl}$ for maxima and minima of total $R_w/c^2$		
	Bow and stern interference component of $R_w/c^2$	Total $R_w/c^2$	Calculated	Measured	
Crest with crest 0.63	0.09	0.85	0.75	0.80	Maximum
Crest with trough 0.49	0.51	0.51	0.51	0.52	Minimum
Crest with crest 0.42	0.44	0.44	0.44	0.44	Maximum
Crest with trough 0.37	0.38	0.38	0.38	0.38	Minimum
Crest with crest 0.34	0.34	0.34	0.34	0.35	Maximum
Crest with trough 0.31	0.31	0.31	0.31	0.31	Minimum
Crest with crest 0.30	0.30	0.30	0.30	0.30	Maximum

these systems, as calculated in the last section of the paper, with the corresponding speeds of maximum and minimum values of  $R_w/c^2$ . This comparison appears in Table II, which also includes for reference the calculated and measured maxima and minima for the shallower model 1302, experiments on this model and the corresponding calculations are described later.

It will be seen that the wave-coincidences agree quite well with the maxima and minima of  $R_w/c^2$ , excepting at the highest speeds where the maxima and minima of  $R_w/c^2$  always occur at a higher speed than that of the wave-coincidences—seriously so in the last maximum. Further, the main discrepancy

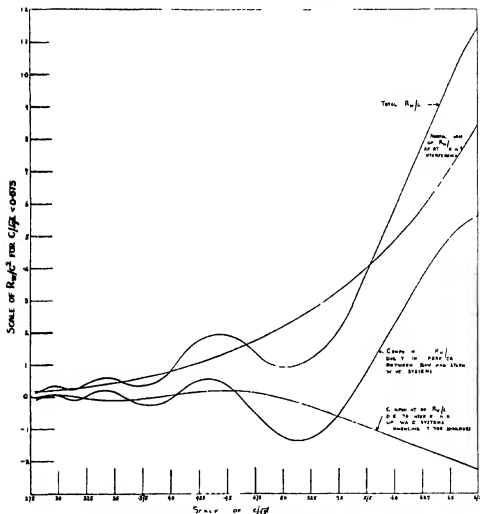


FIG. 5A.—Parabolic model of infinite draft model 1254. Separation of wave resistance into components.  $c/\sqrt{g} < 0.675$

is between the last maximum of total  $R_w/c^2$  and the last maximum of the interference component of  $R_w/c^2$ . This latter difference is due entirely to the term  $1/\kappa^{4/3}$  in the first term in the bracket on the right-hand side of equation (6) this term varies as  $c^4$ , and therefore delays the maximum of the  $R_w/c^2$

curve by causing the function to continue to increase after the interference term has actually begun to decrease. The difference between the highest maximum of the component and the corresponding wave coincidence is due to the factor  $1/\kappa^{2/3}$  outside the bracket, which acts in a similar way. Naturally the effects of these factors are only sensible for large values of  $c$ , and their influence has been checked by separate calculations which confirm the remarks made above. It is noteworthy that the distance from the first crest of the

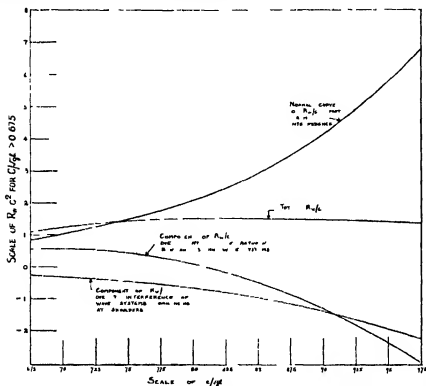


FIG 5B —Parabolic model of infinite draft model 1254 Separation of wave resistance into components  $c/\sqrt{gl} > 0.675$

bow wave to the first crest of the stern-wave, as shown in fig 4, increases from a value equal to the length of the model ( $= 16$  feet) at very slow speeds to a value of just over 20 feet at  $c/\sqrt{gl} = 0.80$ , where  $l$  is, as above, the length of the curved surface of the bow or stern of the form ( $= 7.5$  feet). Owing to the delay in the higher maxima of  $R_w/c^2$ , for reasons which are stated above this distance if deduced from the calculated resistances would appear to increase more than it actually does. Thus Havelock (*loc cit*) has calculated the apparent value of this distance from the resistance of a model, like No 1254 but without

parallel, whose length was 160 feet he finds this distance—equal to  $Z + \lambda/2$  in his notation—to increase from 166 feet at the slowest speeds to 260 feet at  $c/\sqrt{gl} = 0.72$

### *Measurement of Resistance of Shallow Model*

It has been shown that the calculated components of the wave-profiles for this form are related to the maxima and minima of the calculated curves of  $R_w/c^2$ , and that these calculated components add up to form a complete wave-profile which, generally speaking, has the same peculiarities of shape as the wave-profile actually measured with model 1254. It remains to show that the calculated resistance curve agrees in the same way with a measured resistance curve. As explained earlier, the sharp corners at the bilge of model 1254 would render resistance measurements with this model of very doubtful interpretation. Since it is known that draft has very little influence on the shape of resistance curves,\* a model, No. 1302, was constructed having the equation

$$\left. \begin{aligned} y &= b \{1 - \overline{[x - a/l]^2}\} & \text{from } z = 0 \text{ to } z = 1 & \left\{ \begin{array}{l} \text{from } x = l + a \\ \text{to } x = a \end{array} \right. \\ y &= (b/3) \{4 - z^2\} \{1 - \overline{[x - a/l]^2}\} & \text{from } z = 1 \text{ to } z = 2 & \\ y &= b & \text{from } z = 0 \text{ to } z = 1 & \left\{ \begin{array}{l} \text{from } x = a \\ \text{to } x = -a \end{array} \right. \\ y &= (b/3) (4 - z^2) & \text{from } z = 1 \text{ to } z = 2 & \\ y &= b \{1 - \overline{[x + a/l]^2}\} & \text{from } z = 0 \text{ to } z = 1 & \left\{ \begin{array}{l} \text{from } x = -a \\ \text{to } x = a \end{array} \right. \\ y &= (b/3) \{4 - z^2\} \{1 - \overline{[x + a/l]^2}\} & \text{from } z = 1 \text{ to } z = 2 & \end{aligned} \right\} \quad (7)$$

where  $a$ ,  $b$ ,  $l$  have the same values as for model 1254 described above, i.e., 0.5, 0.484, and 7.5 feet respectively. The form of this model is shown in fig. 1, it is the same as model 1254 for 1 foot depth, but then tapers off in a parabolic curve to a knife edge keel at 2 feet depth, the waterlines at any section are similar parabolic arcs joined by 1 foot of parallel body amidships. It will be seen from the equation that there is a slight vertical angle or knuckle at  $z = 1$ , but this was rounded off in the actual model on the sections near amidships, where alone it was sensible. The resistance of this model was measured in the Yarrow tank, using the ordinary resistance dynamometer, over a range of speeds from  $c = 4.0$  to  $c = 12.3$  ft per sec. Since the dynamometer is only capable of measuring 49 lbs. resistance, added pulls of 10 and 20 lbs

\* See Havelock, 'Proc. Roy. Soc., A, vol. 108, pp. 582 et seq. (1925).



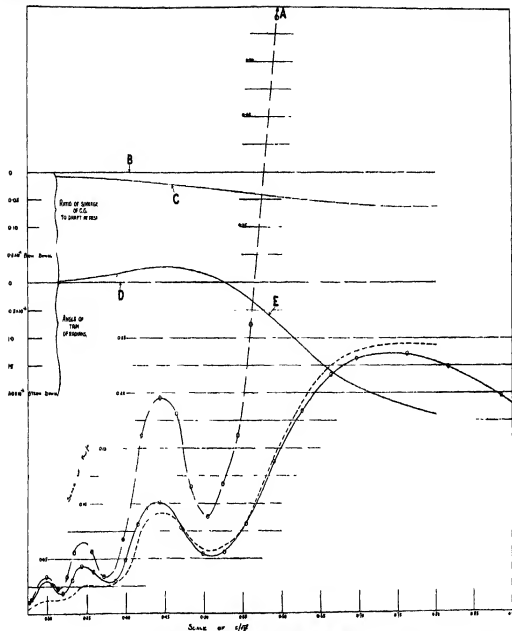


FIG. 6.—Resistance of parabolic models Nos. 1254 and 1202.

A. Curve rises to maximum value of  $R_w/\rho g = 1.0$  at  $c/\sqrt{g} =$  about 0.72. B. Zero line for sinkage curve. C. Curve of sinkage of C.G. (plotted as proportion of draft at rest). D. Zero line of trim curve. E. Curve of angle of trim.

*Key to Resistance Curves.*

- — Calculated curve of  $R_w/\rho g$  for model of infinite draft with vertical sides—approximately Model 1254—load water line same as Model 1202.
- ... Observed curve of  $R_w/\rho g$  for Model 1202. Frictional resistance subtracted as calculated by Froude using surface as actually wetted during motion.
- 0 — Calculated curve of  $R_w/\rho g$  for Model 1202.



were provided as required by cords attached to the model and tensioned by the necessary weights. The speed of advance was measured, as usual, by the aid of an electric contact which opened momentarily at intervals of 20 feet travel, and of a clock recording half-seconds. In addition the trim and bodily sinkage of the model due to its motion were recorded during each experiment, in order to estimate the actual violation of the third assumption mentioned above. The actual measured speeds and resistances are given in Table III.

Table III—Model 1302 Mean Temperature 55.7° Fahrenheit

Speed	Resistance	Speed	Resistance	Speed	Resistance
4.04	3.507	6.71	13.09	9.22	30.11
4.21	3.827	6.87	13.56	9.26	29.26
4.29	4.025	6.96	14.26	9.46	11.94
4.46	4.389	7.08	14.54	9.59	33.00
4.57	4.660	7.21	14.62	9.74	35.83
4.74	5.064	7.32	14.85	9.71	36.49
4.84	5.211	7.47	14.93	9.92	39.37
4.97	5.460	7.54	15.04	10.14	40.18
5.04	5.713	7.67	15.24	10.29	42.01
5.17	6.165	7.81	15.62	10.25	42.69
5.32	6.541	7.93	15.89	10.47	46.44
5.53	7.291	7.98	16.17	10.56	46.59
5.69	7.579	8.18	17.04	10.67	46.82
5.81	7.708	8.23	17.40	10.73	49.25
5.98	8.515	8.43	19.01	11.04	52.15
6.04	8.587	8.46	19.41	11.21	54.94
6.24	9.238	8.70	21.77	11.81	59.65
6.32	10.14	8.79	22.61	12.04	62.71
6.49	11.04	8.93	24.57	12.28	64.74
6.56	11.99	8.96	25.14		

The frictional resistance was calculated from the data of Froude,\* a correction being made when the temperature differed from the standard value of 55° F used by Froude. The area of wetted surface assumed in the calculation exceeded the actual wetted surface of the model when at rest by an amount varying from 1½% at the lowest speed to 5½% at the highest, to allow for the additional surface wetted during motion. This correction was the same as had been measured on a model of similar dimensions used previously. Since the correction is in any case only an approximation, the added error due to this procedure is not of importance.

When the frictional resistance as thus calculated is subtracted from the measured resistance, the measured wave resistance  $R_w$  is found, and in fig 6

\* 'Trans. Inst. Naval Arch.', vol. 29, p. 304 (1888)

is shown the curve of  $R_w/c^3$  as measured for this model. The values of the measured angle of trim (in radians), and of the measured sinkage of the C G of the model (expressed as a fraction of the draft) are also shown in this figure

### *Calculation of Resistance for Shallow Model*

The resistance of this form has also been calculated by the use of the expression found by Michell\* for the wave-resistance of a form obeying the assumptions quoted at the beginning of this paper. For a form having the equation  $y = f(x, z)$  this expression is as follows —

$$\left. \begin{aligned} R_w &= (4\rho g^3/\pi c^3) \int_1^\infty (I^2 + J^2) (\lambda^2/\sqrt{\lambda^2-1}) d\lambda \\ \text{where} \quad I &= \int_0^\infty \int_{-\infty}^{+\infty} f'(x, z) e^{-\lambda^2 g x/c^2} \cos(\lambda g z/c^2) dx dz \\ J &= \int_0^\infty \int_{-\infty}^{+\infty} f'(x, z) e^{-\lambda^2 g x/c^2} \sin(\lambda g z/c^2) dx dz \end{aligned} \right\}, \quad (8)$$

where  $f'(x, z) = \partial \{f(x, z)\}/\partial x$

For model 1302, having the equation (7) above, a simple integration shows that

$$\begin{aligned} J &= -(1024/225 \theta^2) \{\sin \theta - \sin(\theta/16)\} \\ &\quad - (15 - \theta \cos \theta)/16 \{3\psi^3 - 2e^{-\psi} (\psi + 1) + 2e^{-2\psi} (2\psi + 1)\}/3\psi^3 \\ I &= 0, \end{aligned}$$

where

$$\theta = 8\lambda g/c^2, \quad \psi = \lambda^2 g/c^2,$$

and hence

$$R_w/c^3 = 11.98 \int_1^\infty \{F_1(\psi) - F_2(\theta)/\lambda^2 \sqrt{\lambda^2-1}\} d\lambda, \quad (9)$$

where  $\theta$  and  $\psi$  are as above, and

$$\begin{aligned} F_1(\psi) &= \{1 - 2e^{-\psi} (\psi + 1)/3\psi^3 + 2e^{-2\psi} (2\psi + 1)/3\psi^3\}, \\ F_2(\theta) &= \{\sin \theta - \sin \theta/16 - (15 - \theta \cos \theta)/16\}^2/\theta^4 \end{aligned}$$

The functions  $F_1$  and  $F_2$  have been calculated and plotted on a large scale, it is thus a simple matter to calculate the value of the integrand on the right-hand side of the equation (9) for a number of values of  $\lambda$  at each speed. Hence the value of the integral has been obtained by Simpson's rule, care being taken

\* 'Phil Mag,' vol 45, p. 113 (1898)

that a sufficient number of values was used to ensure accuracy in the final result. The apparent infinity at the lower end of the range of integration was avoided by the usual method of assuming that to the order of accuracy required

$$\int_1^{1.001} \{F_1(\psi) - F_2(\theta)/\lambda^3 \sqrt{\lambda^2 - 1}\} d\lambda = \frac{1}{2} \left\{ \left[ \frac{F_1(\psi) - F_2(\theta)}{\lambda^3} \right]_{\lambda=1.001} + \left[ \frac{F_1(\psi) - F_2(\theta)}{\lambda^3} \right]_{\lambda=1.000} \int_1^{1.001} \frac{d\lambda}{\sqrt{\lambda^2 - 1}} \right\}$$

the right-hand side of this equation being easily calculated. At the upper end of the range of integration the approximate integration was at each speed carried out sufficiently far to ensure that  $F_1(\psi)$  had attained to within 2% of its maximum value of unity. The rest of the integral can be expanded in inverse powers of  $\theta_0$ , and the first two terms of the result are given by the equation

$$\text{Correction to integral} = (225\kappa^2/32\theta_0^4) (1 - 2 \sin 2\theta_0/\theta_0),$$

where  $\theta_0$  is the value of  $\theta$  up to which the approximate integration has been carried out.

Table IV - Calculated  $R_w/c^2$  for Model 1302

$c/\sqrt{g l}$	$R_w/c^2$	$c/\sqrt{g l}$	$R_w/c^2$
0.186	0.0039	0.443	0.1013
0.222	0.0086	0.471	0.0788
0.251	0.0085	0.490	0.0540
0.277	0.0105	0.526	0.0561
0.306	0.0265	0.554	0.0820
0.310	0.0178	0.590	0.1379
0.332	0.0303	0.627	0.1840
0.343	0.0435	0.693	0.2164
0.358	0.0377	0.697	0.2315
0.387	0.0303	0.762	0.2355
0.399	0.0490	0.815	0.2234
0.415	0.0816	0.886	0.1974
0.426	0.0954		

The curve of  $R_w/c^2$  as thus calculated is shown in fig. 6 compared with the curve for the same quantity as derived from the model experiments, it will be seen that there is general agreement. The actual calculated values of  $R_w/c^2$  are also given in Table IV. The curve of  $R_w/c^2$  as calculated for model 1254 by the method described above is also shown on the diagram, and it will be seen that apart from amplitude the positions on a speed base of the maxima and minima of all three curves are in good agreement. These are

tabulated in the last two columns of Table II, they have been derived from curves drawn on a much larger scale than can be reproduced here

It appears from Table II and from inspection of all the relative curves in fig 6 that the main features of the wave-making of the shallower three-dimensional model are the same as those of the model of infinite draft, and accordingly must originate in interference of similar wave systems to those relating to that form and described above. As might be expected, the resistance at the higher speeds (corresponding to larger waves) is very much larger for the model of infinite draft, whereas at slower speeds the difference is not so great. It is known that the depth to which the shape of a form affects the wave-making varies, at any rate approximately as the length of the wave generated, and thus is supported by the foregoing observation.

#### *General Conclusions*

The maxima and minima of the resistance curves have been shown to originate in interference between the bow and stern component wave-systems, this being calculated definitely for the model of infinite draft, and inferred from the similarity of the resistance curve for the shallower three-dimensional model. It is noted that the higher speed maxima of the resistance curves tend to occur at higher speeds than would be predicted from the speeds of coincidence of the wave-systems, and the reason for this is given.

This general result is in contrast with that obtained previously from a model\* having sections composed of straight lines meeting at angles at the shoulder, in the latter shoulder wave-making was of primary importance.

It is hoped to make further experiments with models of different shapes, and in this way to obtain some definite relation between form and wave-profile. Particularly it is proposed to investigate a form with hollow water-lines, since straight and convex lines have now been dealt with. Until this has been done it is not wise to draw any general conclusions regarding the wave-making properties of forms not yet investigated, since it is clear that in different forms the relative importance of different portions of the form will vary greatly.

#### *Summary*

The several effects of the various geometrical features of a body on the form of the waves it generates when moving on the surface of water, and on the resistance it experiences, are of theoretical as well as practical interest to

\* See 'Trans N.E. Instn. Eng. Shipb.', vol. 47, p. 153 (1931)

shipbuilders. It is now possible to investigate this influence, for simple forms, by approximate methods of calculation, in this paper an investigation is described for a form having parabolic waterlines, where possible the results of calculation are compared with actual measurements made in the Alfred Yarrow tank at the William Froude Laboratory

It is found that there is a general agreement between measurement and calculation. The main difference between the parabolic forms dealt with here and straight line forms previously investigated is that the ends of the former contribute more largely to the wave resistance

No conclusions can be drawn concerning the wave making effects with other shapes until further experiments and calculations have been made

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### *Study of Electrolytic Dissociation by the Raman Effect II — Nitrates*

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(Communicated by O W Richardson, F R S — Received October 18, 1933)

#### *1 Introduction*

In the first part\* of these investigations by the author, the possibility of studying electrolytic dissociation by the Raman effect has been described with special reference to nitric acid. Accurate quantitative estimation has since† been made of the degree of dissociation in the acid by measuring the intensities of Raman lines corresponding to the dissociated  $\text{NO}_2^-$  ions at various concentrations. The results thus obtained indicated that the dissociation in nitric acid is progressive between the concentrations 1-16 N and not complete as has been assumed for highly dilute solutions.

A comparison of the values of the degree of dissociation obtained from this method with those calculated from conductivity measurements revealed a large discrepancy which was attributed to the uncertainty of the latter. The degree of dissociation cannot be directly evaluated from the values of electrolytic conductivity, which depend upon a number of factors such as viscosity of the solution, mobility of the ions, etc., as there is much uncertainty as to the dependence of conductivity on such factors. With Raman spectra, however,

\* 'Proc Roy Soc.' A, vol 127, p 279 (1930).

† 'Proc. Acad. Sci., Amst.' vol. 33, p. 632 (1930).

the number of molecules or ions is directly proportional to the intensity of the Raman lines corresponding to them. Hence a comparison of the number of molecules or ions at one concentration with that in another is made by simply comparing the intensities of the lines corresponding to them in the Raman spectra taken with the respective concentrations. The accuracy of the method is that attainable in the intensity measurements. Thus we have in this method a reliable means of measuring electrolytic dissociation, and another advantage is its applicability to concentrated solutions, for which at present no theory has been fully developed. In order to determine whether electrolytes are completely dissociated even in concentrated solutions, a systematic study of different electrolytes is being undertaken. The results for nitrates will be described in this communication.

## *2 Raman Spectra of Nitrates in the Crystalline State and in the State of Solution*

The applicability of the Raman effect to the determination of electrolytic dissociation consists in the fact that the undissociated molecules give rise to a set of Raman lines which are in entirely different positions from those arising from the dissociated ions. In nitric acid, for example, it is found that the lines with Raman frequencies equal to 953, 1120, 1306  $\text{cm}^{-1}$  attributed to the undissociated molecules are different from those with  $\delta\nu = 725, 1049$ , and 1357  $\text{cm}^{-1}$  corresponding to the  $\text{NO}_3^-$  ions. The latter set of lines are found to be common to solutions of all nitrates, the former being entirely absent from them. With dilution of the acid the lines with  $\delta\nu = 953, 1120$ , and 1306  $\text{cm}^{-1}$  progressively diminish in intensity, while the other set actually increase in intensity up to a certain dilution and then diminish, a phenomenon similar to the variation of electrolytic conductivity with dilution, which has been explained as being caused by dissociation of the electrolyte.

To determine whether there is a similar change in the Raman spectra with undissociated to dissociated nitrates, such spectra are taken with crystals as well as solutions of these salts. Table I contains the results of measurements of Raman frequencies in the two different states.

There are three generalizations which can be deduced from this table. Firstly, the frequencies in the crystalline state are different for different kations. Secondly, for the same kation, the frequency in the solid state is higher than for the solution, with the exception of ammonium and possibly potassium nitrates. And thirdly, the frequencies of all nitrates in solution are identical.



All the three frequencies given in Table I are well known to be due to inner vibrations of the nitrate radical. Leaving alone the two fainter lines with mean  $\delta\nu = 726$  and  $1390\text{ cm}^{-1}$ , the intense line  $1049\text{ cm}^{-1}$  is attributed to the inactive vibrations of the three oxygen atoms symmetrically with respect to the nitrogen atom. Any change in the above frequency can be brought about only by a change in the distances of the oxygens from the nitrogen atom. This can be accomplished by imposing upon this group a strong electric field. This is perhaps what happens in the crystalline state.

Table I - Raman frequencies in  $\text{cm}^{-1}$  of nitrates in crystals and solutions

Kation	$\nu_1$		$\nu_2$		$\nu_3$	
	Crystal	Solution	Crystal	Solution	Crystal	Solution
Lithium*	728	—	1086	1070	1391	—
Sodium	726 (1)	721 (1)	1071 (10)	1049 (10)	1389 (1)	1381 (1)
Potassium	711 (1)	730 (1)	1051 (10)	1046 (10)	1359 (1)	1357 (1)
Ammonium	709 (1)	728 (0)	1043 (10)	1050 (10)	—	1381 (0)
Magnesium	—	—	1039 (10)	1049 (10)	—	—
Calcium	—	722 (0)	1064 (10)	1049 (10)	—	—
Strontium	—	720 (0)	1054 (10)	1049 (10)	—	1355 (0)
Barium	—	—	1049 (10)	—	—	—
Lead*	—	—	1045	—	—	—

\* The values for the crystalline state for these two substances are taken from Gierlich (Ann Physik, vol 5, p 196 (1930))

X-ray analysis has revealed that different nitrates crystallize in different forms and belong to different systems of symmetry. Thus the relative disposition of the kations and anions is different for different crystals.

Sodium nitrate has hexagonal crystals, which appear to have the rhombohedral symmetry of the calcite group. Its structure may be pictured as a sodium chloride grouping in which the nitrate groups replace the chlorine atoms, and which has been compressed along a trigonal axis normal to the plane of the nitrate groups. Thus the sodium atoms are symmetrical with respect to the nitrate groups and form an ionic lattice, in which no single sodium ion can be exclusively attached to any single nitrate ion. If the nitrogen atom is supposed to occupy the centre of the nitrate group, it can be concluded from the above that the nitrogen and sodium atoms are uniformly distributed in the space lattice. But the plane containing the oxygen atoms is not thus situated. The nitrogen atom is not exactly in the above plane, but is slightly

outside it as is revealed by X-ray study. Thus owing to the electrostatic field arising out of the kations there may be a further displacement of the nitrogen atom with respect to the plane containing the oxygen atoms, which would not have taken place if there was no such field. The change in the nitrate frequencies may be due to this displacement. Since the nitrates with different kations belong to different systems of symmetry the relative disposition of the kations and the anions will be different. Hence the disposition of the resultant electrostatic field with respect to the plane of the oxygen atoms in the nitrate group may be different. Thus the displacement of the nitrogen atom from its mean position, and hence its distance from each of the oxygen atoms, will be slightly different for different crystals, and consequently the internal frequencies will differ.

For crystals belonging to the same type of symmetry, as with calcium, strontium, barium, and lead nitrates, in which the relative disposition of the kations and the anions is the same, there is a gradual diminution in frequency from the lighter to the heavier elements. This regular change may be explained as follows. The heavier elements, having a larger ionic volume, are at greater distances from the anions, and hence the resultant electrostatic field, to which the anion is subject, is smaller in intensity thus resulting in a smaller displacement of the nitrogen atom.

The equality of the Raman frequency in solutions of all nitrates indicates the identity of this radical in all of them. This can occur only if it is present in the completely dissociated condition. If there were undissociated molecules, the kation with which the nitrate group is in combination should give rise to a strong electrostatic field (much stronger than in the ionic lattice of the crystal) involving a change in the distance of the nitrogen with respect to the oxygens. As the different kations of the same valency, which have the same charge, are of different diameters, they should be at different relative distances from the centre of the nitrate group. Hence the displacement of the nitrogen atom from the centre of the group should be different. Hence the Raman frequency also should change from one nitrate to another.

In order to detect whether there are even faint lines in positions different from that of the strong line  $\delta\nu = 1049 \text{ cm}^{-1}$ , very long exposures with solutions of ammonium and sodium nitrates of concentrations 12 N and 8 N respectively were given. Though the above line gave as much density on the photographic plate as some of the strong mercury lines, no trace of any additional line could be seen though the concentrations were nearly the highest possible, thus giving confirmatory evidence of the identical nature of the nitrate

ion in solutions of all nitrates, and consequently of their complete dissociation in solution

If it is assumed that there is complete ionization, the anions and cations are uniformly distributed in space, with the result that there are nearly as many anions surrounding a nitrate group as there are cations. Thus the resultant electrostatic field, if there be one at all, is negligible compared to that present in the crystalline state. Also the nitrate groups, on account of their freedom of movement, are oriented in all possible directions with respect to the neighbouring ions and hence there is no possibility of any displacement of the nitrogen ion with respect to the plane containing the oxygen atoms. Hence the frequency of the nitrate ion is undisturbed by the neighbouring ions.

Other evidence for the hypothesis of complete dissociation is obtained from nitrates containing water of crystallization. Magnesium and strontium nitrates containing respectively six and four molecules of water of crystallization have shown in the solid state lines of very small Raman frequencies which belong to the outer vibrations. Though they are very strong in the crystalline state they are entirely absent in solution. This shows that the complex group containing the six  $H_2O$  molecules and the two  $NO_3$  groups are completely disrupted in solution, so that they are no longer intact to produce the outer vibrations.

### 3 Variation with Concentration of the Raman Line of Sodium Nitrate

On lines similar to the study of dissociation in nitric acid, the variation, with concentration, of the intensity of the Raman line with  $\delta\nu = 1049\text{ cm.}^{-1}$  for sodium nitrate has been investigated. Raman spectra with 6.0, 4.8, 3.6, and 2.4 N concentrations are taken under identical conditions of illumination, temperature, time of exposure, etc., so that any variation in the intensity of the Raman line can only be due to variation in the number of radicals giving rise to it. The intensity of the Raman line at  $4226\text{ \AA}$  due to excitation by the 4047 line of the mercury arc spectrum is given in Table II.

Table II -Intensity of the  $1049\text{ cm.}^{-1}$  Raman line at different concentrations

Concentration (	Intensity $I_s$	$I_s/C$	$I_s/C + 1/8 \text{ at } 0$
6.0 N	1.9	0.65	100
4.8 N	2.9	0.61	94
3.6 N	2.2	0.61	94
2.4 N	1.6	0.67	103

The first column contains the molal concentration, the second the intensity of the Raman line, the third the ratio of the intensity to concentration, and in the fourth column are given the ratios of the values of  $I_s/C$  to the corresponding value for 6.0 N taken as 100

#### 4 Discussion of the Results

The values in the last column are constant to within the degree of accuracy that is attainable in intensity measurements. There is no progressive change and if the slight variation owing to the inaccuracy in the intensity determination is neglected they are fairly constant. This is an indication that the number of  $\text{NO}_3$  ions at any concentration is proportional to the concentration. There is no evidence of any other change taking place as was found for nitric acid, wherein the diminution in intensity is much less than that in proportion to concentration. The above is therefore a clear evidence for the complete dissociation of sodium nitrate even at high concentrations.

The results from conductivity data lead to a different conclusion. In Table III is shown the degree of dissociation calculated from both the formulae

$$\alpha = \frac{\Lambda_s}{\Lambda_0},$$

where  $\Lambda_s$  and  $\Lambda_0$  are the equivalent conductivities at concentrations  $c$  and 0 respectively as well as

$$\alpha = \frac{\Lambda_s \eta_0}{\Lambda_0 \eta_c}$$

where  $\eta_c$  and  $\eta_0$  are the viscosities

Table III - Degree of dissociation in sodium nitrate from conductivity data

Concentration $c$	Equivalent conductivity $\Lambda_s$	$\frac{\Lambda_s}{\Lambda_0}$	Viscosity $\eta$	$\frac{\Lambda_s \eta_0}{\Lambda_0 \eta_c}$
0.0 N	105.33	100	1.00	100
0.1 N	87.24	82.8	1.00	82.8
0.2 N	83.28	78.1	1.01	78.0
0.5 N	74.05	70.3	1.02	71.7
1.0 N	65.80	62.5	1.04	65.0
2.0 N	54.5	51.8	1.15	59.7
3.0 N	48.0	45.7	1.27	55.5
4.0 N	39.2	37.2	1.40	52.1

The values of the degree of dissociation calculated from both the formulae indicate that the dissociation in sodium nitrate progressively increases with increasing dilution.

This discrepancy can be explained by the fact that the Raman effect method is direct involving no uncertainties, the only assumption made is that the intensity of the Raman line is proportional to the number of  $\text{NO}_3^-$  ions giving rise to it, which seems to be quite legitimate. The conductivity formula however, is uncertain as has already been pointed out.

The sources of evidence for the hypothesis of complete dissociation of nitrates are the following -

- (1) All nitrates in solution give rise to exactly the same Raman line, though in the solid state they reveal different lines. This shows that the nature of the radical corresponding to this line is identical in all of them in the state of solution, which cannot be so unless they are completely dissociated.
- (2) Raman spectra even with saturated solutions of nitrates taken with very long exposures did not show any trace of additional lines which can be attributed to undissociated molecules.
- (3) In some nitrates containing water of crystallization, *e.g.*, magnesium and strontium, the external frequencies which are revealed by the solid state are entirely absent in solution. This indicates that the molecular aggregates are completely disrupted in solution.
- (4) Measurements of the variation of intensity of the  $4226 \text{ \AA}$  line of sodium nitrate with concentration have revealed a direct proportionality between the two, this should occur only if the substance is completely dissociated even at high concentrations.

#### Summary

Raman frequencies of nitrates in the crystalline state and in the state of solution are given. While in the former state the frequencies are different for different nitrates, in the latter they are identically the same. This is taken as an indication of the complete dissociation of nitrates in solution. Spectra with large concentrations of sodium and ammonium nitrates, taken with very long exposure, did not reveal even the slightest trace of any additional line due to undissociated molecules. Determination of the variation of intensity of the  $\text{NO}_3^-$  line of sodium nitrate with concentration has shown that the ratio of intensity to concentration is constant, indicating the exact proportionality between number of  $\text{NO}_3^-$  ions and concentration. It is therefore concluded that the study of the Raman effect leads to the hypothesis of complete dissociation of nitrates even at high concentrations.

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*The Theory of the Structure of Ethylene and a Note on the Structure of Ethane* †

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There are two fairly distinct methods of attempting the solution of the structure of polyatomic molecules. One is associated principally with the names Hund and Mulliken, while the other owes its development chiefly to Hentler, London, Slater and Pauling. The former of these, which following Van Vleck, ‡ we shall denote as the H-M procedure, allows the electrons to be fed, one at a time, into a self-consistent field possessing the symmetry of the nuclear framework. Each energy level can absorb two electrons on account of the spin degeneracy, and the levels are filled in succession until all of the electrons have been used up. In contrast, the H-L-S-P method is based on the idea of electron pairs. The spin of an electron on one nucleus is coupled with that of another electron on another nucleus to give a resultant spin or zero, and the pair then form a saturated bond. Pairing occurs between the spins in such a way as to give a maximum bonding energy, and the number of free spins capable of being paired with electrons on other atoms constitutes the valency of an atom. While the two modes of attack, when carried to a complete solution, are equivalent, they often bear little resemblance to each other, to the approximation to which they are generally tractable. It is therefore important, when considering any problem of valency, to make calculations both ways, and if the results agree, one can feel fairly sure of their accuracy. It is the principal aim of the present paper to attempt such an attack on the ethylene molecule  $\text{CH}_2=\text{CH}_2$ . In addition, some interesting results will be given on the ethane molecule  $\text{CH}_3-\text{CH}_3$ .

Pauling§ and Slater|| have suggested rather tentatively that with ethylene the four valence electrons on the C nuclei are arranged with tetrahedral symmetry, just as they are in the methane molecule. This would make the H-C-H angle  $109.5^\circ$ , and the two pairs of electrons comprising the double bond,

† A preliminary account of the earlier portions of this paper were given at the 1933 Washington Meeting of the American Physical Society.

‡ 'J. Chem. Phys.', vol. 1, pp. 177, 219 (1933).

§ 'J. Amer. Chem. Soc.', vol. 63, p. 1367 (1941).

|| 'Phys. Rev.', vol. 37, p. 481 (1931).

similar. In order to get the maximum overlapping, and hence presumably the least energy, all six nuclei must lie in one plane. We shall discuss this model in some detail and show that by a slight redistribution of the bonds an even more stable arrangement is possible. Mulliken† has proposed essentially the same type of double bond as we employ, but he did not make any detailed calculations to establish the point. Hückel‡ has also considered the ethylene molecule, and he concluded that it is the  $\pi$ - $\pi$  bonds which prevent free rotation about the C-C joint. We shall establish this point in much greater detail than he did, and prove at the same time that if one of the  $\text{CH}_2$  groups is rotated, the position of minimum energy is attained when all six nuclei lie in one plane.

There is one position of the nuclei, "the right-angled model," that appears at first sight more stable than any other. If the HCH angles are made right angles and the HCH planes parallel, it is possible to have a pure  $p$ -bond between each H and its C atom, and for the double bond one pure  $s$ - $s$  and one pure  $p$ - $p$  bond. The criterion that the most stable configuration is that with the most overlapping would seem to favour this arrangement. However, a closer examination shows that there are other arrangements which involve a large amount of hybridization,§ sufficient to render them even more stable than the somewhat curious model just described.

One can obtain very simply a rather entertaining result on the ethane molecule. It is that there is free rotation about the C-C joint, both in the H-M and the H-L-P-S approximations, provided interactions between any two hydrogen atoms and between any hydrogen atom and the distant carbon atom, are neglected.

We proceed to give these ideas more exact formulation.

#### THE H-L-P-S APPROXIMATION FOR ETHYLENE.

Our computations are based on the well-known formula of perfect pairing

$$W = Q + \sum_i J_{ii} - \frac{1}{2} \sum_{i \neq k} J_{ik}, \quad (1)$$

where  $W$  is the energy of the configuration apart from atomic terms,  $Q$  is the Coulomb energy, and the  $J$  are defined as the diatomic two-electron integrals

$$J = \iint \psi^*(a_1) \psi^*(b_2) H \psi(b_1) \psi(a_2) dv_1 dv_2 \quad (2)$$

† 'Phys. Rev.,' vol. 43, p. 279 (1933)

‡ 'Z. Physik,' vol. 60, p. 423 (1930)

§ When wave functions  $f$  and  $g$  from two different families have nearly equal energies, they may "hybridize," so that the functions to be taken are  $af + bg$  and  $cf + dg$

Here  $a_i$  represents the co-ordinates of electron  $i$  relative to the nucleus  $a$ , and  $H$  is the Hamiltonian operator. The summation is over all pairs  $(i, k)$ , a  $-\frac{1}{2}J_{ii}$  term arising from any two not coupled to zero spin,  $i = e$ , not forming a bond, and a  $J_{ii}$  from each pair forming a bond. Our calculations bring out very clearly the importance of *hybridized* wave-functions, by allowing some  $2s$  wave-function to be mixed in with the various  $2p$  wave-functions, still retaining the aggregate carbon configuration  $2s(2p)^3$ , the bonds are made very much stronger. In consequence, the molecule constructed with them gains enormously in stability. As is usual, we shall neglect the  $(1s)^2$  shell of carbon, the interaction between any two hydrogen atoms, and the interaction

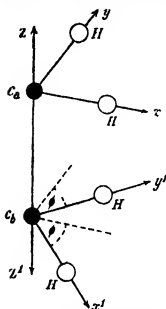


FIG 1 - The figure illustrates the right angled model. The bonds between the C and H atoms are pure  $2p$  and the double bond is made up of one  $(s, s)$  and one  $(p, p)$  bond. The energy of the configuration does not depend on  $\phi$ .

between any hydrogen atom and the distant carbon atom. Furthermore, we shall ignore the presence of the higher quantum orbits of both C and H in the expansion of the complete wave-function in terms of the unperturbed C and H wave-functions.

It is necessary to introduce two kinds of exchange integrals, the  $C$  which have four suffixes, and the  $N$  which have two. These integrals are exhibited in (4). The  $C$  arise from two electrons which are always on one or the other carbon nuclei. Two of the suffixes define the initial state and two the final. With the integrals  $N$  there is always one electron on a hydrogen nucleus and one on a carbon nucleus. Two suffixes are sufficient here to define the integral since it is assumed that an electron on a hydrogen nucleus is always in a  $(1s)$  state.

We consider now several plausible arrangements of the ethylene molecule. Since the "right angle" model is by far the simplest algebraically, we start with it.

*The Right-angle Model*—Let us arrange the model so that the C-C line is vertical, denote the upper C nucleus by  $C_a$  and the lower by  $C_b$ . Take as the axis of  $z$  the upward drawn line through the two carbon nuclei, and as the  $x$  and  $y$  axes, the lines joining  $C_a$  to its two H companions, such that  $(x, y, z)$  is an orthogonal, right-handed system of axes,



of origin  $C_a$ . The lower  $CH_3$  group may be specified by the left-handed system ( $x', y', z'$ ), of origin  $C_b$ , where  $z'$  is taken in the opposite sense to  $z$ . Denote by  $\phi$  the angle between the  $x$  and the  $x'$  axes.

The choice of wave-functions is as follows —

$$\left. \begin{array}{llll} \psi(H_1) & , & \psi_a(2p\sigma_a) & , & \psi(H_2) & , & \psi_a(2p\sigma_a) \\ \psi(H_3) & , & \psi_b(2p\sigma_b) & , & \psi(H_4) & , & \psi_b(2p\sigma_b) \\ \psi_a(2s) & , & \psi_b(2s) & , & \psi_a(2p\sigma_z) & , & \psi_b(2p\sigma_z) \end{array} \right\} \quad (3)$$

where members of a pair with balancing spins are separated by a comma, and pairs are separated by semi-colons. Here  $\psi(H)$  is simply the wave-function of the  $(1s)$  state of  $H$ . By  $\psi_a(2p\sigma_z)$  we mean that this is a  $2p$  wave-function on the  $C_a$  nucleus,  $z$  being the axis of quantization and  $\sigma$  denoting that  $m_l$  is zero with reference to this axis. This wave-function clearly has the form  $\mathcal{H}(r_a)$ . There are, however, two  $\pi$  wave-functions, corresponding to  $m_l = \pm 1$  respectively, and these may be taken in either real or complex form. We prefer the real form and take  $\psi_a(2p\sigma_a) = \psi_a(2p\pi_x)$ , which has the form  $\mathcal{H}(r_a)$ , and  $\psi_a(2p\sigma_b) = \psi_a(2p\pi_y)$  which has the form  $\mathcal{H}(r_a)$ . The equality signs are a consequence of the fact that a  $2p$  wave-function which is of the  $\sigma$  type with reference to one axis of quantization is of the  $\pi$  type when referred to an axis of quantization perpendicular to the former. The notation  $\pi$  and  $\pi'$  is to remind ourselves that there are two types of  $\pi$  wave-functions.

Neglecting the atomic and Coulomb parts of the energy, we obtain on using (2) and (3) in (1) —

$$W = -U_{aaaa} - (C_{aaaa} + C_{vvvv} + 2C_{avav} + 2C_{vvav} + C_{vvvv} + C_{vvvv} - 4N_{aa} + 4N_{vv} + 2N_{av})$$

where

$$\left. \begin{array}{l} C_{avav} = -\frac{1}{2} \iint \{ \psi_a(\alpha_1) \psi_b(\beta_2) H \psi_b(\gamma_1) \psi_a(\delta_2) \\ \quad + \psi_a(\delta_1) \psi_b(\gamma_2) H \psi_b(\beta_1) \psi_a(\alpha_2) \} dv_1 dv_2 \\ N_{av} = -\frac{1}{2} \iint \{ \psi_a(\alpha_1) \psi_b(\alpha_2) H \psi_b(\beta_1) \psi_a(\beta_2) \\ \quad + \psi_a(\alpha_1) \psi_b(\beta_2) H \psi_b(\alpha_1) \psi_a(\beta_2) \} dv_1 dv_2 \\ H/e^2 = \frac{1}{r_{a1}} + \frac{1}{r_{b1}} - \frac{1}{R} - \frac{1}{r_{12}}, H'/e^2 = \frac{1}{r_{a1}} + \frac{1}{r_{b2}} - \frac{1}{R} - \frac{1}{r_{12}} \end{array} \right\} \quad (4)$$

Here, for example,  $\psi_a(\alpha_1)$  means the wave-function of electron 1 on the  $a$  nucleus in a quantum state  $\alpha$ . The minus sign has been inserted into the

definition of the integral  $C$  to make certain of the more important integrals positive. Subscripts  $\pi$  and  $\pi'$  only occur together, and the primes denote that the  $\pi$  and  $\pi'$  wave-functions have their axes of symmetry perpendicular to each other. It may be remarked here that any integral  $C$  with one or three  $\pi$  suffixes, or any integral  $N$  with one  $\pi$  suffix vanishes, as can be seen very simply from the symmetry properties of the wave-functions.

The angle  $\phi$  does not appear in the expression for the energy, and hence within the limits of our approximation there is free rotation about the C-C

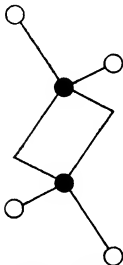


FIG. 2—The figure illustrates an ethylene molecule built on tetrahedral wave functions (Slater and Pauling's original suggestion). This arrangement is not as good as that shown in fig. 3.

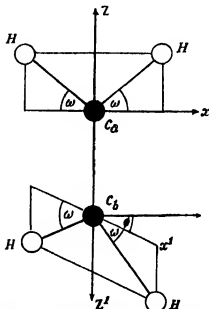


FIG. 3—The figure illustrates the  $\omega$  model. When  $\phi = 0$  we have "plane ethylene" and when  $\phi = \pi/2$ , "perpendicular ethylene". The angles  $\omega$  and  $\phi$  are to be chosen to make the energy a minimum.

joint. We postpone a discussion of the signs and relative magnitudes of the various integrals till later.

*The Slater-Pauling Model*—Next in order of simplicity is the Slater-Pauling model. The reader will probably understand immediately from the figure the way that the various atoms are disposed in this particular arrangement. Each C atom has four tetrahedral valence arms, two of which point away symmetrically from the remote C nucleus, and are saturated with H atoms.

The four remaining valence arms are coplanar and constitute the double bond. It is seen that all six nuclei lie in one plane and that the HCH angle has the tetrahedral value  $109.5^\circ$ . The appropriate carbon wave-functions are

$$(3/4)^{1/2} [\psi(2p, \sigma_i) + (1/3)^{1/2} \psi(2s)],$$

where  $\psi(2p, \sigma_i)$  means that the  $2p$  wave-function is directed along one of the four tetrahedral lines radiating from the C nucleus.

After some manipulation, the energy is found to be

$$\begin{aligned} W = & 5(C_{ssss} + C_{ssss})/16 + C_{ssss}/4 + C_{ssss} + 5(2C_{ssss} + 2C_{ssss} + C_{ssss})/8 \\ & - 3(2C_{ssss} + 2C_{ssss} + C_{ssss})/8 - 3(2C_{ssss} + 2C_{ssss} + C_{ssss})/8 + 3(C_{ssss} \\ & + C_{ssss})/4 + 3(C_{ssss} + C_{ssss} + C_{ssss})/2 - 5N_{ss}/2 + \frac{1}{2}N_{ss} + 4N_{ss} - 3(3)^{1/2}N_{ss} \end{aligned}$$

where the integrals are defined by equation (4). As before, Coulomb and atomic energies have been omitted. We shall return to this formula later.

*The  $\omega$ -Model*—We now discuss the arrangement which we believe to be the one actually realized in nature. It is assumed that the plane of either  $\text{CH}_2$  group contains the other C nucleus. The model then has two angular parameters  $\omega$  and  $\phi$  whose values are to be chosen so that the energy has its minimum value. The choice of directed wave-functions for this model has been given by Van Vleck (*loc cit*). Construct on the nucleus  $C_a$  a right-handed system of axes ( $x, y, z$ ) with  $C_a z$  pointing away from  $C_b$  and with the  $xz$  plane containing  $H_1$  and  $H_2$ . Construct on the  $C_b$  nucleus a similar system of axes ( $x', y', z'$ ), these, however, being left-handed with  $H_3$  and  $H_4$  lying in the  $x'-z'$  plane. Denote by  $\phi$  the angle between the  $x$  and the  $x'$  axes, and by  $\omega$  any of the four equal angles  $H_1 C_a x$ , ( $\pi - H_2 C_a x$ ),  $H_3 C_b x'$ , ( $\pi - H_4 C_b x'$ ).

The directed wave-functions are as follows—

$$\left. \begin{aligned} & \psi(H_1), \quad \psi(H_2), \quad \psi(H_3), \quad \psi(H_4), \\ & \psi_a(2p, \sigma_1), \quad \psi_b(2p, \sigma_{11}), \\ & (1 + \cos 2\omega)^{1/2} [\psi_a(2p, \sigma_1) + (\cos 2\omega)^{1/2} \psi_a(2s)], \quad s = H_1 \text{ and } H_2 \\ & (1 + \cos 2\omega)^{1/2} [\psi_b(2p, \sigma_{11}) + (\cos 2\omega)^{1/2} \psi_b(2s)], \quad s = H_3 \text{ and } H_4 \\ & [k(\omega)]^{1/2} [-\operatorname{cosec} \omega \cos 2\omega \psi_a(2p, \sigma_1) + (\cos 2\omega)^{1/2} \psi_a(2s)], \\ & [k(\omega)]^{1/2} [-\operatorname{cosec} \omega \cos 2\omega \psi_b(2p, \sigma_{11}) + (\cos 2\omega)^{1/2} \psi_b(2s)], \\ & \text{with} \end{aligned} \right\} \quad (5)$$

$$1/k(\omega) = \operatorname{cosec}^2 \omega \cos^2 2\omega + \cos 2\omega$$

Proceeding exactly as before, one can obtain the expression for the energy. After collecting terms and rejecting some which vanish from symmetry properties, we find that

$$\begin{aligned}
 W = & U_{xxxx} (1 - 3 \cos^2 \phi/2) + (U_{xxxx} (1 - 3 \sin^2 \phi/2) + 2(U_{xx\tau\tau} + 2U_{xx\sigma\sigma} + U_{\sigma\sigma\sigma\sigma} \\
 & + \frac{1}{2} (-11 + 12 \sec^2 \omega - 3 \sec^4 \omega) (U_{\sigma\sigma\sigma\tau} + \frac{1}{2} (-2 + 6 \sec^2 \omega - 3 \sec^4 \omega) U_{\sigma\sigma\sigma\sigma} \\
 & - 3 \tan^2 \omega \sec^2 \omega \cos 2\omega (U_{\sigma\sigma\sigma\sigma} + U_{\sigma\sigma\sigma\tau} + U_{\sigma\sigma\sigma\sigma}) \\
 & + 6 \sin \omega \sec^4 \omega (\cos 2\omega)^{\frac{1}{2}} [\cos 2\omega (U_{\sigma\sigma\sigma\sigma} \\
 & + \sin^2 \omega (U_{\sigma\sigma\sigma\tau}) + (2 - 3 \sec^2 \omega) N_{\sigma\sigma} + (-1 + 3 \sec^2 \omega) N_{\sigma\tau} + 4 N_{\sigma\sigma} \\
 & - 6 \sec^2 \omega (\cos 2\omega)^{\frac{1}{2}} N_{\sigma\sigma}
 \end{aligned}$$

As usual the atomic and Coulomb terms are omitted. These are presumably the same for all models: the atomic terms need not concern us, but we give now the expression for the Coulomb energy  $Q$

$$Q = D_{\sigma\sigma\sigma\sigma} + D_{\sigma\sigma\sigma\tau} + 2D_{\sigma\sigma\sigma\sigma} + 2D_{\sigma\sigma\sigma\tau} + 4D_{\sigma\sigma\sigma\sigma}$$

$$+ 4D_{\sigma\sigma\sigma\tau} + 2D_{\sigma\sigma\sigma\sigma} + 4M_{\sigma\sigma} + 4M_{\sigma\tau} + 8M_{\sigma\sigma}$$

where

$$\begin{aligned}
 D_{\sigma\sigma\sigma\sigma} = & \frac{1}{2} \left\{ \left\{ \psi_a(\alpha_1) \psi_b(\beta_2) H \psi_a(\gamma_1) \psi_b(\delta_2) \right. \right. \\
 & \left. \left. + \psi_a(\gamma_1) \psi_b(\delta_2) H \psi_b(\alpha_1) \psi_a(\beta_2) \right\} dv_1 dv_2, \right.
 \end{aligned}$$

$$\begin{aligned}
 M_{\sigma\sigma} = & \frac{1}{2} \left\{ \left\{ \psi_H(\gamma_1) \psi_c(\alpha_2) H' \psi_H(\sigma_1) \psi_c(\beta_2) \right. \right. \\
 & \left. \left. + \psi_H(\sigma_1) \psi_c(\beta_2) H' \psi_H(\sigma_1) \psi_c(\alpha_2) \right\} dv_1 dv_2, \right.
 \end{aligned}$$

and  $H$  is defined by (1). Notice that  $D$  and  $M$ , unlike  $U$ , have no minus sign in their definition.

A brief explanation of why the angle  $\phi$  does not appear in the expression for the energy of the right-angled model while it does in that of the  $\omega$ -model, may be given here. The essential point is that with the right-angled model there are no  $\pi$ - $\pi$  bonds, and when the summation  $-\frac{1}{2} \sum J_{ij}$  for interactions between bonds is made, the dependence of  $W$  on  $\phi$  disappears. With the  $\omega$ -model, however, there is a  $\pi$ - $\pi$  bond, and as the contribution to the energy from a bond has a coefficient  $+1$  (compared with  $-\frac{1}{2}$  for electrons in different bonds), the angle  $\phi$  still remains explicitly in  $W$ .

We have now expressions for the energy of three plausible models. The important question arises as to the signs and magnitudes of the various integrals. We devote a special section to a discussion of this point.

*Discussion of Signs and Magnitudes of Various Integrals*—There are three possible ways of determining the size and approximate magnitude of most of the integrals appearing in our equations. The simplest is the criterion of maximum overlapping. This would tell us, for example, that  $C_{ssss}$  is positive and quite large,  $N_{ss}$  is negative and fairly small, and so on. The second possibility is from the work of Bartlett and Furry† on the  $Li_2$  and  $Be_2$  molecules. We can obtain some very useful information from the tables given in their papers. The third possibility is from the celebrated figure given by Mulliken‡ for the behaviour of the energy levels of various diatomic molecules as the internuclear distance is decreased from large distances to small. On the whole, the three methods agree fairly well, and fortunately, those integrals whose sign is doubtful are small, and not of much consequence.

Bartlett and Furry in their work use conjugate complex wave functions  $(x \pm iy)f(r)$  while we use the real wave-functions  $x f(r)$  and  $y f(r)$ . It is therefore necessary to get the transformation scheme from their integrals to ours. This is obtained without much trouble by the use of simple trigonometric identities

$$\begin{aligned} C_{ssss} &= -I_{10}, \quad C_{ssss} = J'(ss, 00), \quad C_{ssss} = J'(ss, 11) \\ C_{ssss} &= -J'(s1, s1), \quad C_{ssss} = -I_{11}, \quad C_{ssss} = -I_8 \\ C_{ssss} &= -\frac{1}{2}I_8, \quad C_{ssss} = -I_8 - \frac{1}{2}I_8, \quad C_{ssss} = \frac{1}{2}[J'(ss, s0) - J'(ss, 0s)] \end{aligned}$$

The remainder of our integrals do not occur in the expressions for those energy levels which were calculated.

Perhaps the most interesting integrals are  $C_{ssss}$  and  $C_{ssss}$ , since it is the relative magnitude of these two which determines whether the angle  $\phi$  is zero or  $\pi/2$ , i.e., whether "plane" or "perpendicular" ethylene is the more stable. According to Bartlett and Furry  $I_8$  is small and positive, while  $I_2$  is much larger in magnitude and negative in sign. As a consequence, the  $\pi$ - $\pi$  bonds are bonding. This agrees with Mulliken's figure. The criterion of maximum overlapping does not give very definite information on the sign and magnitude of these two integrals, but our conclusions seem very reasonable when this test is applied to them. We therefore have obtained the important result that plane ethylene is more stable than perpendicular. If  $C_{ssss}$  were negative and larger in absolute magnitude than  $C_{ssss}$ , the  $\pi$ - $\pi$  bonds would

† J. H. Bartlett, jr., 'Phys. Rev.', vol. 37, p. 507 (1931), Furry and J. H. Bartlett, jr. *ibid.*, vol. 38, p. 1615 (1931), vol. 39, p. 210 (1932).

‡ 'Rev. Mod. Phys.', vol. 4, p. 40 (1932).

be anti-bonding if coupled to zero spin, but then, of course, we would couple them to a spin of unity. For the dependence of  $W$  on  $\phi$  we then find

$$C_{xxxx}(1 + \frac{1}{2} \cos^2 \phi) + C_{xxzz}(1 + \frac{1}{2} \sin^2 \phi),$$

and once again the plane model is the more stable. The fact that ethylene is not paramagnetic rules this case out.

One can substitute the numerical value of  $C_{xxxx}$  as calculated by Bartlett (*loc cit*), to evaluate the dependence of the energy  $W$  on  $\phi$ . It is found that  $W = 3I_z \cos 2\phi/4$ . Taking an effective nuclear charge of unity and an internuclear distance of 8 gives 0.176 volts as the total variation with  $\phi$ . For  $R = 6$ , the figure is 0.376 volts. These values seem rather low, especially as the latter is, in a way, an upper limit. Dr Sutherland has kindly informed the writer that from data (not yet published) on the infra-red spectrum of ethylene, he has concluded that there is a fundamental twisting frequency of about 750 wave-numbers (1/10 volt). Eucken and Parts† have observed independently a similar value from their work on specific heat determinations. When allowance is made for the half-quantum zero point energy, it is seen that a lower limit for the depth of the trough is about 1/6 volt ‡.

According to overlapping considerations  $C_{xxxx}$  is large and positive, and one would expect it to be the greatest of all the integrals. Mulliken (*loc cit*) has it large and positive, but Bartlett has it negative. This gives him  $(\sigma\sigma)^2 \Sigma_u^-$  for the ground state of a molecule formed from two similar atoms, each with a single  $2p$  electron, an incorrect result. The reason has been elucidated by Stehn,§ who concluded that the Heitler-London first order perturbation theory would give very poor results for any state involving a  $\sigma_u$  electron. There seems little doubt therefore, that  $C_{xxxx}$  is positive and large.

The integral  $C_{zzzz}$  is positive by all three methods, and in magnitude is fairly large. One would expect all of the integrals  $C_{xxzz}$ ,  $C_{yyzz}$ , and  $C_{zzzz}$  to be positive, and of about equal magnitude, somewhere between that of  $C_{xxxx}$ .

† 'Z. Phys. Chem.,' B, vol. 20, p. 184 (1933).

‡ [Note added in proof, January 10, 1934.—Applying the results of Koenig on eigen values in periodic fields of potential ('Phys. Rev.' vol. 44, p. 657, 1933), we find that in order to account for a fundamental twisting frequency of 750  $\text{cm}^{-1}$ , it is necessary to have  $C_{xxxx} = 0.7 \text{ v.e.}$  This gives about 1 v.e. for the difference of energy between plane and perpendicular ethylene. According to Pauling and Wheland ('Jour. Chem. Phys.,' vol. 1, p. 362, 1933),  $C_{xxxx}$  is about  $1\frac{1}{2} \text{ v.e.}$  Chemical evidence on the heat of activation of various compounds derived from ethylene seems to favour the former of these estimates. The calculations will be described in a note to the Physical Society of London.]

§ 'Phys. Rev.,' vol. 42, p. 582 (1932).

and that of  $C_{ss}$ . Now the choice of relative phase of the  $2s$  and  $2p$  wave-functions is still at our disposal and we can therefore take  $C_{ss\sigma}$  and  $C_{ss\pi}$  as negative. Since  $N_{s\sigma}$  is of a type similar to  $-C_{ss\sigma}$ ,  $N_{s\sigma}$  will be positive, a circumstance very favourable to the  $\omega$ -model. Summing up, we have approximately

$$C_{ss\sigma\sigma} > -C_{ss\pi\pi} > C_{ss\sigma\pi} = C_{ss\pi\sigma} = C_{ss\pi\pi} > -C_{ss\sigma\sigma} > C_{ss\pi\pi} > 0,$$

where the equality signs are not to be taken too seriously, but are probably satisfied fairly well.

We consider next those integrals with two  $\pi$ -subscripts. Both  $C_{ss\pi\pi}$  and  $C_{ss\pi\sigma}$  are almost certainly negative and small, but the remainder seem very uncertain. However, they are not capable of affecting our results to any extent and so we do not trouble about them. There still remain the integrals involving two H and two C wave-functions. These are well known (Van Vleck, *loc cit*),  $N_{ss}$  and  $N_{ss}$  are positive and  $N_{ss}$  negative. We have already chosen  $N_{s\sigma}$  positive

$$N_{s\sigma} > N_{ss} > 0 > N_{ss}$$

*Comparison of Energies of Different Models* --Using the results of the previous sections, we can make a critical comparison of the different models. For the  $\omega$ -model there are two parameters  $\omega$  and  $\phi$ , but we shall, of course, take  $\phi = 0$  throughout, for reasons already explained. The best value of  $\omega$  is probably a small one, perhaps about  $25^\circ$ . To facilitate comparison with the Slater-Pauling model, however, let us take for  $\omega$  the tetrahedral value  $\sin \omega = (1/3)^{1/2}$ . By doing this the hydrogen energies  $N$  in the  $\omega$ -model are made exactly equal to those in the Slater-Pauling model, and the comparison is simplified accordingly. We find for the difference in energy

$$W = -[3(C_{ss\sigma\sigma} + C_{ss\pi\pi})/16 - 3(C_{ss\sigma\sigma} + C_{ss\pi\pi})/4 + 3(C_{ss\sigma\pi} + C_{ss\pi\sigma} + C_{ss\pi\pi})/4 + 3C_{ss\pi\pi}/4] + 3[(C_{ss\sigma\sigma} + C_{ss\pi\pi} + C_{ss\sigma\pi} + C_{ss\pi\sigma} + C_{ss\pi\pi} + C_{ss\pi\pi}) - 2(C_{ss\sigma\sigma} + C_{ss\pi\pi} + C_{ss\pi\pi})]/4$$

The terms in the first bracket all favour the  $\omega$ -model. The terms in the second bracket are doubtful, but in any case nearly cancel each other out. We conclude that even with this value of  $\omega$ , the  $\omega$ -model is more stable than the Slater-Pauling model. A few words may well be spent in explaining what the difference really is between the Slater-Pauling model and the  $\omega$ -model with  $\phi = 0$  and  $\omega$  the tetrahedral value. The difference is solely in the double bond. The  $\omega$ -model has a strong ( $s, \sigma$ ) bond along the C-C axis and a weak

( $\pi$ - $\pi$ ) bond perpendicular to this axis. The Slater-Pauling model has two equivalent bonds which are not very strong because they are too much inclined to the C-C axis.

Let us now compare the  $\omega$ -model with the right angle model. For convenience take  $\omega = 30^\circ$ . We find

$$W_{\omega=30} - W_r = [5C_{\sigma\sigma\sigma\sigma}/6 + 4(C_{\pi\pi\pi\pi}/3 + 2(2)^{\frac{1}{2}}(2C_{\sigma\sigma\pi\pi} + C_{\pi\pi\sigma\sigma})/3 - 3C_{\pi\pi\pi\pi}/2 - 2(C_{\sigma\sigma\pi\pi} + C_{\pi\pi\sigma\sigma} + C_{\pi\pi\sigma\sigma})/3] + [2(N_{\sigma\sigma} - N_{\pi\pi}) - 4(2)^{\frac{1}{2}}N_{\sigma\pi}]$$

There does not seem any doubt that this quantity is negative, so that the  $\omega$ -model is the more stable of the two. An even more convincing proof can be obtained by taking, for example,  $\cot \omega = 3$ , but the expressions for the energy are rather too long to print. It should be noticed that it is the hybrid terms which gain the day for the  $\omega$  model.

#### THE H-M APPROXIMATION FOR ETHYLENE

It is possible to make calculations on the ethylene molecule by the H-M method. The secular determinants for the various models are readily set up, the difficulty is to solve them. The assumptions that we shall make and the general procedure that we shall adopt are very similar to those described by Van Vleck (*loc cit*). We give only the barest minimum description of these and refer the reader to Van Vleck's paper for further details. A certain number of interesting results can be obtained without much trouble, but others need a very complete examination before their validity is established. The trouble is briefly this: the magnitudes of the differences of the binding energies of carbon and hydrogen as they appear in the perturbation problem are only about equal to those of some of the resonance integrals. If, therefore, one tackles the problem by perturbation theory, there are difficult questions of convergence, and on this account one may obtain quite the wrong result. In addition, there is the further complication that not even the order of the roots is known for certain. There are in all twelve electrons to be established in the molecule, so that allowing for spin-degeneracy, the lowest six of the twelve roots of the secular determinant are required. Two are undoubtedly much lower than the rest, and two higher, but the order of the remaining eight, of which four are needed, is somewhat uncertain. According as to which four are considered lowest, so the degree of the "ionic-ness" of the resulting molecule is varied. If one assumes, for example, what is certainly not so, that those four of the eight are much the lowest which are built up on carbon root approximations, then the molecule is very ionic and the majority of the



charge is around the two carbon nuclei, leaving the hydrogen nuclei very bare. All one can do in a situation such as this is to consider different variations in the order of the roots which may correspond to physical reality, to make plausible assumptions for the relative magnitudes of the resonance integrals and to examine how the stability of the molecule changes as these various factors are changed. By such a process we attempt to show that plane ethylene is the most stable.

The same general method of attack is to be adopted as was employed in the H-L P-S approximation, that is, we shall endeavour to discover which of several possible models is the most stable. From symmetry arguments, it is very reasonable to suppose that the plane of one  $\text{CH}_2$  group and the plane of the other  $\text{CH}_2$  group are either (a) coincident, or (b) perpendicular, or (c) parallel, and that the two hydrogens in either  $\text{CH}_2$  group are equivalent and symmetrically situated with respect to the C atom. We could decide between (a) and (b) by minimizing the energy with respect to the azimuth  $\phi$ , and, in fact, we set up the secular determinant in a way that would permit this. It turns out, however, that the determinant is so much simplified by putting  $\phi = 0$  or  $\phi = \pi/2$ , that it is worth our while to make these substitutions directly and compare the roots in the two cases. To get the best results from (c) when it is agreed that H-H repulsions are to be neglected, one would naturally take the HCH angle as  $\pi/2$ . In other words, (c) is the analogy of our old 'right-angle' model. We now wish to prove that (a) is better than (c). This must be accomplished by a suitable choice of the angle  $\omega = (\pi - \text{HCH})/2$ , the best value of which we believe to be small. It should be noticed that there is no point in introducing a case (d), based on tetrahedral wave-functions, because this would only give us the same secular determinant in another guise, and would change neither the roots nor the final wave-functions.

The molecular wave-functions for the ethylene molecule are to be taken as linear combinations of the twelve wave-functions

$$\psi(1s, \text{H}_i), \quad (i = 1, 2, 3, 4)$$

$$\psi(2s, \text{C}_j), \quad \psi(2p_{\sigma}, \text{C}_j), \quad \psi(2p_{\pi}, \text{C}_j), \quad \psi(2p_{\pi}, \text{C}_j), \quad (j = 1, 2)$$

suitably centred around their appropriate nuclei. The axes of quantization ( $x, y, z$ ) may be conveniently taken as an orthogonal set with the  $z$  axis corresponding with the C-C line. These wave-functions are solutions of the equations

$$[\nabla^2 + (8\pi^2m/\hbar^2)(W_k^0 - V_C)] \psi(2k, \text{C}) = 0 \quad (k = s \text{ or } p),$$

$$[\nabla^2 + (8\pi^2m/\hbar^2)(W^H - V_H)] \psi(1s, \text{H}) = 0,$$

where  $V_H$  and  $V_C$  are suitable "self-consistent field" potential energies of an electron in the field of an H and C atom respectively. For the convenience of the reader, we redefine Van Vleck's integrals, which have one suffix or none, and some of our own, which have two —

$$\left. \begin{aligned}
 B &= \int \psi(1s, H_i)^2 V_C dv, & C_k &= - \int \psi(2k_i, C)^2 V_H^i dv, \\
 D_H &= - \int \psi(2s, C) \psi(2\sigma_i, C) V_H^i dv, \\
 Q_k &= - \int \psi(1s, H_i) \psi(2k_i, C) V_H^i dv, \\
 R_k &= - \int \psi(1s, H_i) \psi(2k_i, C) V_C dv, \\
 T_k &= \int \psi(1s, H_i) \psi(2k_i, C) dv, \\
 M_{ka} &= - \int \psi_a(k_a, C)^2 V_C^b dv, & S_{kl} &= \int \psi_a(k_b, C) \psi_b(l_a, C) dv, \\
 D_C &= - \int \psi_a(2s, C) \psi_a(2\sigma_b, C) V_C^b dv, \\
 N_{kl} &= - \int \psi_a(k_b, C) \psi_b(k_a, C) \{V_C^a + V_C^b\} dv
 \end{aligned} \right\} \quad (b)$$

Here  $k$  and  $l$  represent  $\pi, \sigma$ , or  $\nu$ . The suffix on  $k$  or  $l$  denotes the axis of spatial quantization. Once again  $a$  and  $b$  refer to the carbon nucleus, and  $i$  to the hydrogen nucleus, which is under consideration. Notice that  $T$  and  $S$  are non-orthogonality parameters whereas the rest of the integrals are energy parameters.

*The  $\omega$ -model* — We shall now set up the secular determinant for the  $\omega$ -model. There are two possible ways of obtaining it. Either very simple wave-functions may be employed and the determinant so obtained reduced by the application of some elementary transformations, or suitable wave-functions may be taken to give the required determinant immediately. The former is the natural line of attack, but on account of the difficulties of printing we adopt the latter method of approach, thereby avoiding the necessity of printing the secular determinants more than once.

Let us adopt a system of axes similar to that we used before, a right-handed set  $(x, y, z)$  on the upper nucleus  $C_a$  and a left-handed set  $(x', y', z')$  on the lower nucleus  $C_b$ . The  $x$ -axis and the  $x'$ -axis are taken parallel, and

$z$  and  $z'$  are taken collinear but of opposite sense. We denote by  $\omega$  any of the four equal angles  $(\pi - \text{HCH})/2$ , and by  $\phi$  the azimuth of  $H_1$  as measured from  $C_{\infty}x$ . Then the azimuth of  $H_2$  is naturally  $(\pi + \phi)$ . The azimuth of  $H_3$  we maintain as zero, and the azimuth of  $H_4$  as  $\pi$ . Consider the following twelve wave-functions, arranged in two sets of six --

- |  |  |
|--|--|
| (1) $[\psi_a(2s) + \psi_b(2s)]/(2)^{1/2}$ ,                  | (6) $[\psi_a(2s) - \psi_b(2s)]/(2)^{1/2}$ ,                  |
| (2) $[\psi_a(2p\sigma_z) + \psi_b(2p\sigma_z')]/(2)^{1/2}$ , | (5) $[\psi_a(2p\sigma_z) - \psi_b(2p\sigma_z')]/(2)^{1/2}$ , |
| (3) $[\psi(H_1) + \psi(H_2) + \psi(H_3) + \psi(H_4)]/2$ ,    | (4) $[\psi(H_1) + \psi(H_2) - \psi(H_3) - \psi(H_4)]/2$ ,    |
| (7) $[\psi_a(2p\sigma_x) + \psi_b(2p\sigma_x)]/(2)^{1/2}$ ,  | (12) $[\psi_a(2p\sigma_x) - \psi_b(2p\sigma_x)]/(2)^{1/2}$ , |
| (8) $[\psi_a(2p\sigma_y) + \psi_b(2p\sigma_y)]/(2)^{1/2}$ ,  | (11) $[\psi_a(2p\sigma_y) - \psi_b(2p\sigma_y)]/(2)^{1/2}$ , |
| (9) $[\psi(H_1) - \psi(H_2)]/(2)^{1/2}$ ,                    | (10) $[\psi(H_3) - \psi(H_4)]/(2)^{1/2}$                     |

On calculating the matrix elements  $\int \psi_\alpha H \psi_\beta$ , where  $\alpha$  represents any one of the upper six and  $\beta$  any one of the lower six wave-functions,  $H$  being the Hamiltonian, it is found that the integral is zero. In other words, this choice of wave-functions has factorized the secular equation of order twelve into two each of order six. Moreover, the first of these sixth order equations again factors into two cubics. It is convenient, in the arrangement of these equations, to number the rows and columns to correspond with the way the wave-functions are numbered. The two  $(s, \sigma)$  cubics may be written together in the form

$F_{\pm}$	$a$	$b$
$a$	$G_{\pm}$	$c$
$b$	$c$	$H$

where

$$F_{\pm} = W_s^c - 2C_s \pm N_{ss} - M_{ss} \pm S_{ss}W - W,$$

$$G_{\pm} = W_p^c - 2(\sin^2 \omega C_p + \cos^2 \omega C_r) \pm N_{pp} - M_{pp} \pm S_{pp}W - W,$$

$$H = W^H + B - W, \quad a = N_{ss} - S_{ss}W - 2\sin \omega D_H + D_s,$$

$$b/(2)^{1/2} = -Q_s - R_s + T_sW, \quad c/(2)^{1/2} = -\sin \omega (Q_p + R_p - T_pW)$$

The sixth order  $(\pi, \pi)$  equation does not factorize with a general value of  $\phi$ . It is

$$\begin{array}{c|cc|cc|cc} J & p & s & r & p & q \\ \hline p & K & t & O & -q & p \\ \hline & t & H & O & t & s \\ \hline & O & O & H & O & r \\ \hline p & -q & t & O & K_1 & p \\ \hline q & p & s & r & p & J \end{array} \quad (7)$$

where

$$J_1 = W_p^2 - \cos^2 \omega (1 + \cos^2 \phi) (C_\sigma - (1 + \sin^2 \omega - \cos^2 \omega \cos^2 \phi) C_\pi) \\ \pm N_{\pi\pi} - M_{\pi\pi} \pm S_{\pi\pi} W - W,$$

$$K_1 = W_p^2 - \cos^2 \omega \sin^2 \phi C_\sigma - (2 - \cos^2 \omega \sin^2 \phi) C_\pi \\ \pm N_{\pi\pi} - M_{\pi\pi} \pm S_{\pi\pi} W - W,$$

$$H = W^H + B - W, \quad p = \cos^2 \omega \cos \phi \sin \phi (C_\sigma - C_\pi)$$

$$q = \cos^2 \omega \sin^2 \phi (C_\sigma - C_\pi) \quad l = Q_\sigma + R_\sigma - T_\sigma W,$$

$$r = -l \cos \omega \quad s = r \cos \phi, \quad t = r \sin \phi$$

Two results are immediately clear. In the first place, there is a decided hybridization of  $s$  and  $\sigma$  wave-functions, but no hybridization of  $s$  and  $\pi$  or of  $\sigma$  and  $\pi$  wave-functions. While this does not agree with the H-L-S-P viewpoint, represented by (5), there is no cause for alarm, since it is the type of contrast continually arising between the two methods. The second result is, that as far as the dependence of the energy on  $\phi$  is concerned, we can confine our attention to the second of these determinants, since  $\phi$  does not occur in the first. We do have to decide, however, which roots must be taken out of the one determinant and which out of the other.

Let us for the present confine our attention to the second of these determinants. If we put  $\phi = 0$ , two of the roots, viz.,  $K_1$ , separate out completely, and one easily reduces the remaining quartic to two quadratics, which can be solved exactly

$$\begin{array}{c|c} J_1 - r(2)^{\frac{1}{2}} & \\ \hline r(2)^{\frac{1}{2}} & H \end{array} \quad \begin{array}{c|c} J_1 - r(2)^{\frac{1}{2}} & \\ \hline r(2)^{\frac{1}{2}} & H \end{array} \quad \text{with } \phi = 0$$

If we put  $\phi = \pi/2$  in the determinant, there is again considerable simplification and it factorizes immediately into two identical cubics

$$\begin{array}{ccc|c} J_- & r & q & \\ \hline r & H & -r & \\ \hline q & -r & J_+ & \end{array} \quad \text{with } \phi = \pi/2, J_+ = K_+.$$

The question of the determination of the roots of this cubic now arises. One is tempted to use perturbation theory, but unfortunately the results by this method are not satisfactory. In the first place the ordinary formulæ must be modified to allow for the presence of the off-diagonal elements in  $W$ . Fortunately, these are already of second order, a circumstance which allows the roots to be obtained to the fourth order without too much extra complication. One can assume that the carbon roots are a good way below the hydrogen, and thus expand in powers of  $r/(J - H)$ , etc., taking for the four required roots those built on  $J_-, K_-, J_+, K_+$ . Alternatively, one can assume that the  $H$  roots are much lower than the  $C$ , and again the expansion can be made in powers of  $r/(J - H)$ . This time, however, only the two  $H$  roots are required from this determinant, and from the  $(s, \sigma)$  determinant are to be taken the other two  $H$  roots and the lowest two  $C$  roots. With either approximation, one finds that it is not until the fourth order terms are included that there is any difference between  $\phi = 0$  and  $\phi = \pi/2$ . Even then, the fourth order terms practically balance out, leaving a residual of equivocal sign. Although this is rather a troublesome state of affairs, it is really due to our having limited the size of the various parameters sufficiently to render the perturbation approximation valid. The difference between the energies of plane and perpendicular ethylene can hardly be calculated as a fourth order term, although it may be rather small.

As a last resource, it was decided to try numerical values in the equations and to examine the rotational stability in this way. The roots are not very difficult to obtain numerically once the values of the various quantities have been specified, and a simple addition of roots decides immediately between  $\phi = 0$  and  $\phi = \pi/2$ . We have tried wide ranges of values in the different parameters, and it seems probable that with values corresponding roughly to those in the actual molecule  $\phi = 0$  is the best arrangement. However, the rather amusing result is obtained that for a highly ionic molecule perpendicular

ethylene is very slightly more stable than plane ethylene. We give now a few illustrative examples. For the angle  $\omega$  we assume a value  $30^\circ$ , for the non-orthogonality parameters  $T_{\sigma} = 0.25$ ,  $S_{\sigma\sigma} = 0.20$ , and for the various energy parameters, in volts

$$H - W_p^0 + M_{\sigma\sigma} = 1, \quad N_{\sigma\sigma} = 3, \quad Q_{\sigma} + R_{\sigma} = 2(3)^{\dagger}, \quad C_{\sigma} = \frac{1}{2}, \quad C_{\pi} = -1/6, \\ N_{\pi\pi} \text{ and } N_{\sigma\pi} \text{ equal and each 5 volts.}$$

Later calculations must decide whether these values are anywhere near the truth or not. More important for our present purpose are the ratios of the various energies, and these should not be too inaccurate. The roots are to be taken as follows: two C and one H from the  $(s, \sigma)$  determinants and one C and two H from the  $(\pi, \pi)$  determinants. It is found that with this very non-polar molecule, plane ethylene is at least  $1\frac{1}{2}$  volts better than perpendicular ethylene (we cannot give the exact figures without specifying the  $(s, \sigma)$  determinants exactly). This is almost certainly too large a difference  $\dagger$ .

If we take  $H + M_{\sigma\sigma} - W_p^0 = 10$  volts, and leave all the other quantities the same, we get a very ionic molecule with most of the charge around the C nuclei. This time perpendicular ethylene is about  $1/25$  volt better than plane ethylene. If we take  $H + M_{\sigma\sigma} - W_p^0 = 3$ , and reduce  $N_{\sigma\sigma}$  to 1 volt, we still have a fairly ionic molecule, and it turns out that perpendicular ethylene is  $1/16$  volt better than plane. With  $H + M_{\sigma\sigma} - W_p^0 = 2$  and  $N_{\sigma\sigma} = 2$ , perpendicular ethylene is  $0.28$  volt more stable than plane. By increasing  $N_{\sigma\sigma}$  or decreasing  $H + M_{\sigma\sigma} - W_p^0$  beyond these values, however, plane ethylene soon becomes more stable than perpendicular. This is caused by the root  $K_+$  of the plane configuration becoming positive, and an H root for it can then be taken from the  $(s, \sigma)$  determinant. The above calculations can be very easily modified where the hydrogen roots are below the carbon, and one obtains the same conclusions as before, that for a molecule which is not too highly ionic, plane ethylene is more stable than perpendicular. While it cannot be said that our calculations by the H-M method decide definitely in favour of  $\phi = 0$ , they do at least reveal how great will be the difficulties of making exact computations of the energy difference between plane and perpendicular ethylene by this method. One would certainly expect the variation of the energy with  $\phi$  to be very much more than in the very symmetrical ethane molecule, where it is only the H-H repulsions that give anything of total amount  $1/60$  volt  $\ddagger$ . Probably the true situation is somewhere

$\dagger$  Perhaps not after all. See footnote, p. 174.

$\ddagger$  Eyring, 'J. Amer. Chem. Soc.', vol. 54, p. 319 (1932).

between our numerical examples one and four, perhaps a little nearer to one. If this is so, then plane ethylene is better by a fraction of a volt, but it is not fair to claim that we have proved this.

We have now to examine the dependence of the energy on  $\omega$ . This problem also is too complicated for even an approximate solution, but one can see in a rough sort of way that a fairly small positive value is required. The one term in our approximation that decides that  $\omega$  is to be positive is  $c$ , just as in the H-L-P-S approximation it was the analogous terms  $C_{\text{axis}}$  and  $C_{\text{over}}$ . The  $(s, \sigma)$  determinants have terms in  $\sin \omega$ , and the roots would have their best values with  $\omega = \pi/2$ , but the  $(\pi, \pi)$  determinants have terms only in  $\cos \omega$  and the roots here would like to have  $\omega = 0$ . Since the diagonal elements in the former determinant are more widely separated than those in the latter, and since the former has only one root strongly affected by  $\omega$  while the latter have two, it is plausible that the best value of  $\omega$  is a small one. It is not obvious at first sight why  $\omega$  is not actually zero, but numerical calculations again elucidate the point and show that  $\omega$  may be small but is not zero.

We have finally to consider the stability of the  $\omega$ -model compared with that of the right-angled model. Once again the situation is such that until some more information is available on the magnitudes of the various quantities appearing in the equations no satisfactory answer can be given. The author was not successful in discovering a transformation which completely eliminated the appearance of  $\phi$  from the secular equation of the right-angled model, but the dependence of the roots on  $\phi$  is obviously less than it is in the  $\omega$ -model. Little is lost, therefore, by putting  $\phi = 0$  directly, and the equation then factorizes into two quadratics and two quartics. While it seems likely that the roots are not as favourable as those of the  $\omega$ -model, the argument is so incomplete that we do not give it. The conclusions of the H-L-P-S approximation on this point must be accepted.

#### THE ETHANE MOLECULE

Calculations very similar to those made on ethylene can be made on ethane. It is a relatively simple matter to write down expressions for the energy, but there is no advantage in doing so, since there can be no doubt of the approximate structure of the molecule, and that is all our calculations can give. There is, however, one interesting result which we shall obtain, namely, that both in the H-L-P-S and the H-M methods of approximation there is free rotation of the

$\text{CH}_3$  groups around the C-C axis, provided interactions of any H atom with the distant C and H atoms are neglected

The choice of directed wave-functions suited to this example is that given by Van Vleck (*loc cit*) for a pyramidal model of methane. We arrange three H atoms around each C atom, symmetrically with regard to the C-C axis. The angle HCC, which we denote by  $\theta$ , and the angle  $\phi$ , by which we denote the azimuth around the C-C axis of one  $\text{CH}_3$  group with respect to the other, then specify the ethane molecule, apart from scale factors. The wave-function of an electron on a C nucleus, entering into a bond with an electron on the other C nucleus, is given by

$$\psi_C = (1 - 2 \cotan^2 \theta)^{\frac{1}{2}} \psi(2p, \sigma_C) - 2^{\frac{1}{2}} \cotan \theta \psi(2s),$$

while any of the three electrons on the C nucleus forming bonds with the electrons of three adjoining H atoms, is given by

$$\psi_H = (2 \operatorname{cosec}^2 \theta/3)^{\frac{1}{2}} \psi(2p, \sigma_H) + (1/3 - 2 \cotan^2 \theta/3)^{\frac{1}{2}} \psi(2s),$$

where  $\sigma_C$  and  $\sigma_H$  mean that the electrons are  $\sigma$  electrons when viewed along the C-C and the C-H lines respectively

We do not attempt to calculate the energy and then minimize it with respect to  $\theta$ . The principal integral, however, is probably  $C_{ssss}$  and one would therefore make its coefficient rather larger than it is with tetrahedral wave functions. In other words, the angle  $\theta$  is probably smaller than the tetrahedral angle and the three C-H lines radiating from each C nucleus must enclose a slightly larger solid angle than they do in methane. The effect of the H-H steric repulsions will be to increase this angle still further.

Let us now consider how the angle  $\phi$  appears in the expression for the energy. Clearly, the energy must have trigonal symmetry expressed by the relation

$$W(\phi) = W(\phi + 2\pi/3) = W(\phi + 4\pi/3)$$

In the expression for  $W$  by the H-L-S-P approximation, the angle  $\phi$  can appear only in a limited number of ways. It can conceivably arise as  $\cos^2 \phi$ ,  $\sin^2 \phi$ ,  $\cos \phi \sin \phi$ ,  $\cos \phi$  and  $\sin \phi$ , each term with different coefficients. But with each term must arise two others obtained by increasing  $\phi$  by  $2\pi/3$  and  $4\pi/3$  respectively. Performing the simple addition, it is easily verified that  $W$  is independent of  $\phi$ . That is, there is free rotation of the  $\text{CH}_3$  groups around the C-C axis, to the approximation we have considered. When one allows for the H-H interactions,  $\phi$  will appear implicitly in the H-H exchange integrals,



because the distance between an H atom of one  $\text{CH}_2$  group and an H atom of the other depends on  $\phi$

We consider now the situation as viewed from the H-M viewpoint. The molecule as defined above by the angles  $\theta$  and  $\phi$  has sufficient symmetry to induce considerable factorization of the secular determinant. There are two types of resonance terms, those between a carbon atom and the hydrogen members of its group, and those between two carbon atoms. We agree to neglect resonance terms between any two hydrogen atoms and between any hydrogen atom and the distant carbon atom. Let us take the C-C axis vertical and choose a right-handed system of axes ( $x, y, z$ ) about the upper nucleus  $\text{C}_a$ , and a left-handed system ( $x', y', z'$ ) about the other nucleus  $\text{C}_b$ . We make  $x'$  parallel to  $x$ ,  $y'$  parallel to  $y$ , and  $z'$  pointing in the opposite direction to  $z$ . The choice of carbon wave-functions which practically diagonalizes the carbon-carbon interaction is

$$\left. \begin{aligned} \psi_a(2s) \pm \psi_b(2s) \\ \psi_a(2p, \sigma_a) \pm \psi_b(2p, \sigma_a) \end{aligned} \right\} \quad (8)$$

$$\left. \begin{aligned} \psi_a(2p, \sigma_a) \pm \psi_b(2p, \sigma_a) \\ \psi_a(2p, \sigma_y) \pm \psi_b(2p, \sigma_y) \end{aligned} \right\} \quad (9)$$

We have now to define the location of the hydrogen atoms. Suppose the lower three to be kept fixed, one of them lying in the  $x'z'$  plane. We can write the wave-functions for these as

$$\psi_b(\text{H}, 0) \quad \psi_b(\text{H}, 2\pi/3) \quad \psi_b(\text{H}, 4\pi/3)$$

Similarly, if we denote the azimuth of one of the three upper H-atoms by  $\phi$ , the wave-functions of these can be written

$$\psi_a(\text{H}, \phi), \quad \psi_a(\text{H}, \phi + 2\pi/3), \quad \psi_a(\text{H}, \phi + 4\pi/3)$$

We are now in a position to write down the secular determinant but it will have such a form that its independence of  $\phi$  is by no means obvious. We introduce the trigonal wave-functions

$$\Psi_1 = [\psi_a(\text{H}, \phi) + \psi_a(\text{H}, \phi + 2\pi/3) + \psi_a(\text{H}, \phi + 4\pi/3)]/(3)^{1/2},$$

$$\Psi_2 = [2\psi_a(\text{H}, \phi) - \psi_a(\text{H}, \phi + 2\pi/3) - \psi_a(\text{H}, \phi + 4\pi/3)]/(6)^{1/2},$$

$$\Psi_3 = [\psi_a(\text{H}, \phi + 2\pi/3) - \psi_a(\text{H}, \phi + 4\pi/3)]/(2)^{1/2},$$

with similar definitions for  $\Psi_4$ ,  $\Psi_5$ , and  $\Psi_6$ , which refer, of course, to the H-atoms on the  $C_2$  nucleus. We now take the combinations

$$\Psi_1, \quad \Psi_6 \quad (10)$$

$$\left. \begin{array}{cc} \cos \phi \Psi_2 - \sin \phi \Psi_3, & \sin \phi \Psi_2 + \cos \phi \Psi_3 \\ \Psi_4, & \Psi_5 \end{array} \right\} \quad (11)$$

It is easily verified that the secular determinant is independent of  $\phi$ , and furthermore that it factorizes directly into a determinant of order six, involving wave-functions (8) and (10), and one of order eight involving wave-functions (9) and (11). Additional factorization is obvious, and one obtains without any difficulty the twice repeated quadratics and the cubics

$$\begin{array}{c} \begin{array}{|cc|} \hline P_+ & \alpha \\ \hline \alpha & H \\ \hline \end{array} \quad \begin{array}{|ccc|} \hline Q_+ & \beta & \gamma \\ \hline \beta & R_+ & \delta \\ \hline \gamma & \delta & H \\ \hline \end{array} \end{array}$$

$$P_+ = W_p^0 + N_{ss} - M_{ss} - 3 \cos^2 \omega (C_s - C_s)/2 - 3C_s - W \pm WS_{ss},$$

$$\alpha = (3)^{1/2} \cos \omega (Q_s + R_s - T_s W), \quad H = W_{11} + B - W,$$

$$Q_+ = W_p^0 \pm N_{ss} - M_{ss} - 3C_s - W \pm WS_{ss},$$

$$R_+ = W_p^0 \pm N_{ss} - M_{ss} - 3(\sin^2 \omega C_s + \cos^2 \omega C_s) - W \pm WS_{ss},$$

$$\beta = -N_{ss} + S_{ss} W + 3 \sin \omega D_H - D_s, \quad \gamma = -(3)^{1/2} (Q_s + R_s - T_s W)$$

$$\delta = (3)^{1/2} (Q_s + R_s - T_s W)$$

We do not attempt the solution of these equations, but it is instructive to exhibit their form. We see that there is considerable hybridization of the  $2s$  and the  $2p\sigma$  wave-functions, but that the initial  $2p\pi$  wave-functions do not acquire any  $2s$  wave-function in the approximation we have considered.

If we insert terms to allow for the interactions between an H and a distant C atom, the secular determinant is much more complicated. Either correction alone is sufficient to spoil the dependence on  $\phi$ , and one has to develop the roots to a high perturbation approximation before this type of integral appears with a coefficient depending on  $\phi$ . In the H-L P-S approximation there is, of course, no dependence on  $\phi$  at all, and it is only the H-H interactions which give anything. Eyring (*loc cit*) has considered this problem and shown that

the most stable position is  $\phi = 0$ , although the fluctuation in energy as  $\phi$  is varied amounts to only 1/60 volts (0.4 kg cal per mol)

Calculations are now in progress on the  $N_2O_4$  molecule to determine which is the most stable one of the three models that have been proposed † Experimental evidence is largely in favour of the one similar in shape to the ethylene  $\omega$ -model discussed at some length in the present paper, but it is as well to have the additional check of theoretical computations. An effort is also being made to extend the calculations of Hückel‡ on the benzene ring, by taking account of the bonds in the plane of the ring as well as those perpendicular to it

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#### SUMMARY

Calculations have been made to determine the structure of the ethylene molecule. Two modes of attack are employed, that of electron pairs and that of molecular orbitals. The two methods give consistent results. It is shown that the most stable arrangement has all six nuclei coplanar, with a large HCH angle, perhaps about  $130^\circ$ . The energy needed to rotate one of the  $CH_2$  groups through  $\pi/2$  about the C-C axis is quite small, probably about  $\frac{1}{2}$  volt.

Similar calculations on the ethane molecule show that it is only the H-H repulsions that prevent the free rotation of the  $CH_3$  groups about the C-C axis.

† Mellor, "Treatise on Inorganic Chemistry," vol. 8, p. 546.

‡ E. Hückel, 'Z. Physik,' vol. 70, p. 204 (1930), vol. 72, p. 310 (1931).

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*Free Paths and Transport Phenomena in Gases and the Quantum Theory of Collisions II The Determination of the Laws of Force between Atoms and Molecules*

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The quantum theory has provided a means of calculating the interaction energies of two atoms by a perturbation method. It appears that the short range interaction forces are due mainly to electron exchange phenomena between the two atoms, while the van der Waals forces arise from mutual polarization effects\*. The theory gives the first of these forces in the first approximation, while the van der Waals forces appear only in the second approximation. At large distances, where the interaction is small, it is somewhat surprising that the first approximation is not sufficient, and one is led to doubt the accuracy of the method when applied at distances at which the first and second approximations give comparable results. At these distances the mutual potential energy is comparable with the mean kinetic energy of a gas atom at ordinary temperatures, and it is therefore clear that a study of gas-kinetic collision phenomena should provide a satisfactory test of the validity of the perturbation method in this region. It is the object of this paper to carry out a number of calculations with this aim in view.

In a previous paper† the quantum theory of collisions was applied to gas-kinetic collisions, and it was shown that, although the classical theory can be used with accuracy to determine the law of force from viscosity and diffusion phenomena associated with heavy gases, it cannot be applied with safety to hydrogen and helium. The method to be used in such cases was given, and it was also shown that the existence of a definite total collision area—a feature of the quantum theory of scattering by a centre of force, the potential of which falls off more rapidly than  $r^{-2}$  at large distances—provides a further means of determining the law of force. As this collision area can now be directly measured with accuracy by molecular ray experiments, the range of applica-

\* J. E. Lennard-Jones, 'Proc. Phys. Soc.', vol. 43, p. 461 (1931).

† Paper I, 'Proc. Roy. Soc.', A, vol. 141, p. 434 (1933).

ability of this method is considerably greater than that of methods based on transport phenomena

In this paper we first consider the interaction of two helium atoms. Using the interaction energy given by Slater,\* the viscosity of helium is computed using the method described in paper I, and it is found by comparison with experiment that although Slater's field is fairly satisfactory, it is not completely accurate. The modifications necessary to improve it are indicated, and the possibility of further test by measurement of free paths is then discussed. Next, the interaction of a helium atom and a helium ion is considered by calculating the mobility of helium ions in helium, taking account of electron transfer processes. In this case it is possible to devise a method of calculating the interaction energy which should be quite accurate, but it is found that the final value for the mobility, calculated with this field, is not in agreement with experiment. The reason for this discrepancy is not clear. Finally, the calculation of free paths for various laws of force between gas atoms is carried out in order to provide material for molecular ray investigation of the interactions.

### §1 The Interaction of Two Helium Atoms and the Viscosity of Helium

The interaction of two helium atoms has been calculated by Slater (*loc cit*), who obtains

$$\{7.7e^{-2.43\rho} - 0.68\rho^{-6}\} \times 10^{-10} \text{ erg}, \quad (1)$$

where  $\rho$  is the nuclear separation in units of  $a_0$ . Margenau† has given a second expression

$$\{7.7e^{-2.43\rho} - 0.68\rho^{-6}(1 + 7.9\rho^{-2} + 30\rho^{-4})\} \times 10^{-10} \text{ erg}, \quad (2)$$

which takes account of dipole-quadrupole and quadrupole-quadrupole interaction terms. These two forms are illustrated in fig. 1.

The viscosity of helium is given by

$$\eta = \frac{5}{3} \frac{M^2}{M^2} \left( \frac{2\pi}{M} \right)^{3/2} \frac{1 + \epsilon}{\pi R_{11}} \quad (3)$$

where

$$J = 1/2 \kappa T, \quad (4)$$

$M$  is the mass of a gas atom, and  $\kappa$  is Boltzmann's constant. The quantity  $R_{11}$  is defined by

$$R_{11} = \frac{1}{2} \int_{-\infty}^{\infty} v^2 Q_1(v) e^{-1/2 v^2} dv \quad (5)$$

\* 'Phys. Rev.', vol. 32, p. 349 (1928)

† 'Phys. Rev.', vol. 38, p. 747 (1931)

where  $Q_v$  is the viscosity cross-section, given by

$$Q_v = 2\pi \int_0^\pi I(\theta) \sin^2 \theta d\theta \quad (6)$$

The angular distribution function  $I(\theta)$  has been defined in paper I, and is such that if a stream of  $N$  atoms per unit area per second is incident on an atom

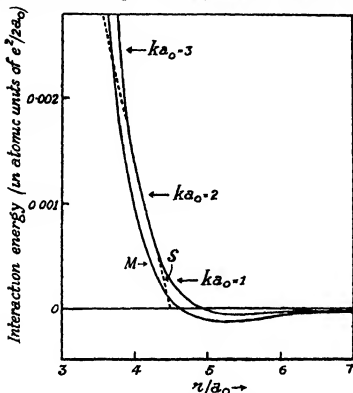


FIG. 1.—Calculated Interaction Energy of Two Helium Atoms S—that due to Slater-Kirkwood. M—that due to Margenau-Slater ——— form of the field

$$O\left(\frac{1}{r^3} - \frac{1}{r_0^3}\right), \quad r < r_0$$

supposed held at rest, the number of incident particles deflected per second between the angles  $\theta$  and  $\theta + d\theta$  is

$$2\pi N I(\theta) \sin \theta d\theta \quad (7)$$

$I(\theta)$  is given, on the quantum theory of collisions, by the expression\*

$$I(\theta) = \frac{1}{2k^2} \left| \sum_n (4n+1) (e^{2ik\delta_n} - 1) P_{2n}(\cos \theta) \right|^2, \quad (8)$$

\* Allowance has been made for the identity of the colliding particles

where

$$k = \pi M v / h \quad (9)$$

corresponding to the velocity  $v$  and the reduced mass  $M/2$  of a pair of colliding atoms, each of mass  $M$ . The phases  $\delta_n$  are obtained from the asymptotic form of the solution of

$$\frac{d^2 u}{dr^2} + \left\{ k^2 - \frac{8\pi^2 M}{h^2} V(r) - \frac{n(n+1)}{r^2} \right\} u = 0, \quad (10)$$

which is zero at the origin, this asymptotic form being

$$u \sim \sin(kr - \frac{1}{2}n\pi + \delta_n) \quad (11)$$

It was shown in paper I how  $\delta_n$  may be calculated if  $V$  is known, by using in the appropriate region methods of approximation due to Jeffreys and to Born for the solution of the equation (10). If we apply this method to the potential (1), we obtain the values, Table I, for  $Q_n$  as a function of the velocity

Table I—Viscosity Cross-sections for Helium, using Slater's Field

$ku_0$	8/9	2/3	4/3	2	4	6
Viscosity cross-section (in units of $a_0^2$ )	46.8	46.0	45.3	41.7	37.5	32.0

Using these values in the appropriate formulae, viz (3) and (5), one finally obtains the values given in Table II for the viscosity of helium at different temperatures—they are compared with the experimental values

Table II—Viscosity of Helium at Various Temperatures

Absolute temperature	Experimental value (in micropoise)	Quantum theory (Slater's field)
294.5	199.4	213
203.1	155.4	165
88.8	91.8	96
20.2	35.03	43
15.0	29.46	36

It will be seen that Slater's field gives values for the viscosity which are about 6% too large at the higher temperatures, while the calculated values are unlikely to be in error by more than 3%. At lower temperatures, however,

the discrepancy is more marked, being of the order of 22 %. This indicates too small a range for the interaction of the two atoms, as the viscosity varies inversely as the viscosity cross-section. Better agreement would be obtained by diminishing the attractive part of Slater's field. Referring to fig 1, we see that Margenau's field is likely to give even less correct values for the viscosity at low temperatures, a result which is confirmed by a detailed calculation \*. These results are to be expected, since the quantum theory includes the possibility of some penetration of the incident waves into the region excluded on the classical theory. At low temperatures it is just possible that some of the discrepancy may be due to the use of the Maxwell-Boltzmann statistics instead of the Bose-Einstein statistics in forming the general equations for viscosity †

In view of the interest of these results, it is important to obtain an independent check on the accuracy of the values of  $Q$ , calculated by the method used. To do this, the following method of calculating the phases  $\delta_n$  was also used for certain chosen values of  $k$ . Suppose we write

$$V = V_0 + v_1, \quad (12)$$

where  $v_1$  is small compared with  $V_0$ . Then equation (10) may be written

$$\frac{d^2(ru)}{dr^2} + \left\{ k^2 - \frac{8\pi^2 M}{h^2} V_0 - \frac{n(n+1)}{r^2} \right\} (ru) - \frac{8\pi^2 M}{h^2} v_1 ru \quad (13)$$

Let us suppose  $V_0$  is chosen so that the equation

$$\frac{d^2(ru_0)}{dr^2} + \left\{ k^2 - \frac{8\pi^2 M}{h^2} V_0 - \frac{n(n+1)}{r^2} \right\} (ru_0) = 0 \quad (14)$$

can be solved analytically to obtain two independent solutions, one of which is zero at the origin and has the asymptotic form

$$ru_0 \sim a_n \sin(kr - \frac{1}{2}n\pi + \eta_n) \quad (15)$$

Since  $u_0$  must represent the sum of an incident plane wave and a scattered spherical wave, we have

$$a_n = e^{i\eta_n} (2n+1) k^{-1} \quad (16)$$

\* It would seem that the above results are in contradiction to the results of paper 1, where the quantum mechanics of the hard sphere model was seen to give larger cross sections than the classical theory. This model has special properties, however, in view of the fact that the wave function of the incident atom must vanish at the boundary of the struck atom, giving a maximum density outside the boundary. For a non-rigid particle, there is some penetration of the incident waves inside the boundary, and the wave function is nearly at a maximum at the boundary.

† Uehling and Uhlenbeck, 'Phys. Rev.', vol. 43, p. 552 (1933)



We may now obtain a further approximation by substituting this solution for  $ru_0$  on the right-hand side of (13), and solving the resultant equation. We then obtain\* for the asymptotic form of the function  $ru$

$$ru \sim a_n \sin(kr - \frac{1}{2}n\pi + \eta_n) + e^{i\epsilon} (2n+1) e^{ikr} \int_0^\infty \frac{8\pi^2 M}{k^3 h^2 a_n^2} v_1 r u_0^2 dr$$

$$\simeq k^{-1} (2n+1) [e^{i\epsilon} \sin(kr - \frac{1}{2}n\pi) + \frac{1}{2}i (1 - e^{2i(\eta_n + \sigma_n)}) e^{ikr}], \quad (17)$$

where

$$\sigma_n = - \int_0^\infty \frac{8\pi^2 M}{k h^2 a_n^2} v_1 r^2 u_0^2 dr, \quad (18)$$

since  $\sigma_n$  must be small compared with unity. The phase  $\delta_n$  will be given by

$$\delta_n = \eta_n + \sigma_n \quad (19)$$

Applying this method to the interaction  $V$ , the form adopted for the potential  $V_0$  was

$$V_0 = C \left( \frac{1}{r^2} - \frac{1}{r_0^2} \right), \quad r < r_0,$$

$$= 0, \quad r > r_0, \quad (20)$$

with the constants  $C$  and  $r_0$  chosen to make the phase  $\sigma_n$  quite small. With this form for  $V_0$  the equation (14) has the solution

$$ru_0 = D_n J_{n+\frac{1}{2}}(\lambda r) + E_n J_{-n-\frac{1}{2}}(\lambda r), \quad r < r_0,$$

$$= A_n J_{n+\frac{1}{2}}(kr) + B_n J_{-n-\frac{1}{2}}(kr), \quad r > r_0, \quad (21)$$

where

$$\lambda^2 = k^2 + \frac{8\pi^2 M}{h^2} \frac{C}{r_0^2},$$

$$v(v+1) = n(n+1) + \frac{8\pi^2 M}{h^2} C \quad (22)$$

Since  $ru_0$  must vanish at the origin,  $E$  must be zero. The ratio of the constants  $A, B, D$  is then determined by the conditions at the boundary  $r = r_0$ , namely, that  $u$  and its derivative be continuous across the boundary. Then  $\eta_n$  is given by

$$\eta_n = \arctan (-)^{n+1} \left( \frac{B_n}{A_n} \right)$$

With appropriate choice of the velocity  $v$  for the calculation, it was possible to arrange that the numerical values of the Bessel functions could be obtained from the very meagre tables available. Having obtained  $\eta_n$  in this way,  $\sigma_n$

\* Massey, 'Proc Roy Soc,' A, vol. 137, p. 447 (1932)

was then calculated from the formula (22) by numerical integration. The form of  $V_0$  used is shown by the broken curve in fig. 1, and it was found that the phase shift  $\sigma$  produced by the difference  $v$  of the Slater field from the field  $V_0$  was seldom more than 0.1 radian. In Table III the values obtained by this method for  $ka_0 = 2$  using the Slater field are compared with those obtained using the methods of approximation due to Jeffreys and Born. It is seen that when the phase is greater than unity, it is given by Jeffreys's method with considerable accuracy. Phases less than, say, 0.1 will be given quite accurately by Born's method. There will generally be only one even phase lying in the intermediate region, and it can be obtained quite accurately by careful interpolation.

Table III—Phases for  $ka_0 = 2$ 

	Accurate calculation	Jeffreys's approximation	Born's approximation.
$\delta_0$	7.63	7.63	—
$\delta_2$	4.90	4.90	—
$\delta_4$	2.70	2.72	—
$\delta_6$	1.17	1.08	—
$\delta_8$	0.27	0.07	1.6
$\delta_{10}$	-0.07	-0.25	-0.07

As the evidence from viscosity indicates a certain inaccuracy in Slater's field, it would be of interest to check these conclusions by measurement of the free paths of helium atoms by molecular ray methods. Therefore, the value of the total cross-section  $Q$  for the Slater field was calculated for various velocities of impact from the formula

$$Q = \int_0^\pi I(\theta) \sin \theta d\theta$$

$$= 8\pi k^{-2} \sum_n (4n+1) \sin^2 \delta_{2n} \quad (23)$$

The values obtained are shown in fig. 2 for various values of  $ka_0$  and the corresponding effective temperatures. It will be seen that definite maxima and minima appear in the curve,\* their occurrence is in no way dependent on the form of the field of interaction, as we saw in paper I that they occurred in the hard sphere model. The effect is due to the presence of a small number of terms only in the series (23) at low velocities, and the fact that only the even

\* This is not to be confused with the ordinary Ramsauer effect, which cannot occur with a repulsive field.

phases appear owing to the identity of the colliding particles. Oscillations of this nature do not occur in the effective cross-section  $Q_e$  for viscosity, for the formula for  $Q_e$  consists of two series of terms of opposite sign, so that such oscillations occur together and tend to cancel out. It will be difficult to confirm the occurrence of the oscillations shown in fig 2, owing to the necessity of using homogeneous beams of atoms, but the marked increase in the cross-section at low temperatures should be observed. It will be noticed that the

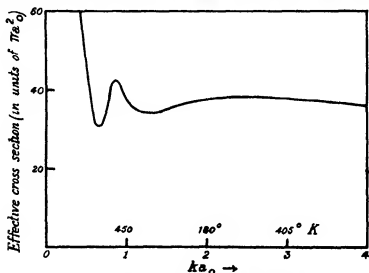


FIG. 2—Effective cross section for the collision of two helium atoms

collision radii corresponding to the cross-section shown in fig 2 are nearly  $\sqrt{2}$  times the classical closest distance of approach, showing that the result obtained for the rigid sphere model in paper 1 holds also for this type of field.

## § 2. The Motion of Helium Ions in Helium. The Mobility and the Interaction $\text{He}^+ - \text{He}$

The mobility of ions in a gas is given by

$$k = eD/kT, \quad (24)$$

where  $e$  is the charge on the ion, and  $D$  is the coefficient of diffusion of the ions in the gas. This coefficient is given by

$$D = \frac{1}{2} \pi^{1/2} \left( \frac{M_1 + M_2}{j M_1 M_2} \right)^{1/2} \frac{1}{v_1 P_{12}} \frac{1}{1 - \epsilon_0}, \quad (25)$$

where  $M_1$ ,  $M_2$  are the masses of the ion and atom respectively,  $v_1$  the number of atoms per cubic centimetre, and  $P_{12}$  is given in terms of a diffusion cross-section  $Q_D$  by the formula

$$P_{12} = 2 \int_{-\infty}^{\infty} V^2 Q_D \exp \left\{ - \frac{2M_1 M_2}{M_1 + M_2} V^2 \right\} dV \quad (26)$$

$Q_D$  is defined in terms of the angular distribution  $I(\theta)$  for collisions between an ion and an atom by

$$Q_D = 2\pi \int_0^\pi I(\theta) \sin^2 \frac{1}{2} \theta \sin \theta d\theta. \quad (27)$$

To calculate  $I(\theta)$  requires a knowledge of the interaction energy between  $\text{He}^+$  and  $\text{He}$ , and, further, we must take into account the possibility of charge transfer from the ion on impact. We will consider the interaction first.

The quantum theory of the interaction of  $\text{He}^+$  with  $\text{He}$  introduces the feature of level splitting due to nuclear symmetry. A helium ion and a helium atom, both in their ground states at infinity, can react in two different ways according as the wave function representing the combined system is symmetric or anti-symmetric in the nuclear co-ordinates. The first of these possibilities gives rise to a potential energy which has a deep minimum (2.5 volts) at a nuclear separation of  $2.1a_0$ , and which only changes to a repulsion at still smaller distances, while the second gives rise to a repulsive interaction, which changes sign only at large distances as a result of the effect of the polarization interaction. The first approximation in the theory, which neglects the polarization, therefore gives rise to two potential energy functions  $v_+(r)$ ,  $v_-(r)$ . Making the usual approximation, we then take for the complete interaction at large distances for the two cases

$$\begin{aligned} V_+(r) &= v_+(r) - C/r^4, \\ V_-(r) &= v_-(r) - C/r^4 \end{aligned} \quad (28)$$

The theory necessary for the evaluation of  $v_+$ ,  $v_-$  has been given by Majorana,\* and by Pauling,† who also calculated the potential energy out to a distance  $4a_0$ . At this distance the energy is of the order of 0.3 volt, showing that the calculation must be extended to much larger distances. This was first done by extending the range of the above-mentioned calculations. It is important to use unperturbed wave functions for the helium atomic electrons which are accurate

\* 'Nuovo Cimento,' vol. 8, p. 22 (1931)

† 'J. Chem. Phys.' vol. 1, p. 56 (1933)

at the large distances involved. Rosen,\* in discussing the technique of this type of calculation, gives the wave function

$$u = 3.20 e^{-1.48r/a_0}, \quad (29)$$

where  $r$  is measured in  $\text{\AA}$ , as accurately representing the charge density of helium at these distances. This wave function was used whenever the overlap of charge densities at large nuclear separations was involved in the integrals concerned. The constant  $C$ , given from the known polarizability of the helium atom, has the value  $2.34 \times 10^{-44}$  in cgs units. The resulting form of the potential functions  $V_s$ ,  $V_a$  is shown in fig 3

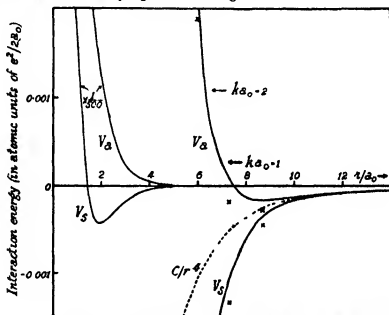


FIG 3 —Calculated interaction energies of a helium ion and a helium atom. Full line curves calculated by method of Majorana Pauling. Crosses give points calculated by the method of Hylleraas.

We must now consider the possibility of charge transfer on collision. After a collision has taken place between a helium atom and a helium ion, it is impossible to distinguish in any way between a scattered ion and a struck atom which has lost an electron, and account of this must be taken in the quantum theory of the collision. This theory has already been discussed by Massey and Smith,† and it is found that the function  $I(\theta)$  must take the form

$$I(\theta) = \frac{1}{2} |f_s(\theta) + f_a(\theta) + f_s(\pi - \theta) - f_a(\pi - \theta)|^2 \quad (30)$$

\* 'Phys. Rev.', vol. 38, p. 255 (1931)

† 'Proc. Roy. Soc. A', vol. 142, p. 142 (1933).

Here  $f_+(\theta)$ ,  $f_-(\theta)$  are the functions

$$\begin{aligned} \frac{1}{2\pi k} \sum_n (e^{2i\beta_n} - 1) (2n+1) P_n(\cos \theta), \\ \frac{1}{2\pi k} \sum_n (e^{2i\gamma_n} - 1) (2n+1) P_n(\cos \theta), \end{aligned} \quad (31)$$

where the  $\beta_n$ ,  $\gamma_n$  are the phase shifts produced by the fields  $V_+$ ,  $V_-$  respectively. Since

$$P_n(\cos \theta) = (-)^n P_n(\cos \pi - \theta),$$

we see that  $I(\theta)$  given by (30) has the same form as for the collision of two neutral atoms, viz.,

$$\frac{1}{2} |f(\theta) + f(\pi - \theta)|^2,$$

with

$$f(\theta) = \frac{1}{2\pi k} \sum_n (e^{2i\delta_n} - 1) (2n+1) P_n(\cos \theta),$$

where the even phases  $\delta_{2n}$  are calculated for the field  $V_+$ , and the odd phases  $\delta_{2n+1}$  for the field  $V_-$ . Then we have, from equation (26) of paper I,

$$\begin{aligned} Q_D &= 2\pi k^{-2} \sum_n \{ (2n+1) \sin^2 \delta_n - 2(n+1) \cos(\delta_n - \delta_{n+1}) \sin \delta_n \sin \delta_{n+1} \}, \\ &= 2\pi k^{-2} \sum_n \{ (2n+1) \sin^2 \delta_n - 2(n+1) (\frac{1}{2} \sin 2\delta_n \sin 2\delta_{n+1} \\ &\quad + \sin^2 \delta_n \sin^2 \delta_{n+1}) \}, \end{aligned} \quad (32)$$

where  $\delta_n = \beta_n$  for  $n$  even, and  $\delta_n = \gamma_n$  for  $n$  odd. The calculation of the phases proceeds in just the same way as in § 1, but we are now concerned with a large number of terms owing to the big range of the potentials  $V_+$ ,  $V_-$  and the calculation of  $Q_D$  must be simplified in the following way.

The series (33) is broken up into two parts, thus

$$Q_D = Q_D^1 + Q_D^2,$$

where

$$\begin{aligned} Q_D^1 &= 2\pi k^{-2} \sum_{n=0}^m, \\ Q_D^2 &= 2\pi k^{-2} \sum_{n=-m}^{\infty}. \end{aligned} \quad (33)$$

For a particular value of  $k$ , the value of  $m$  is so chosen that  $\beta_n$  rapidly becomes very large as  $n$  decreases to values less than  $m$ , while  $\gamma_n$  also becomes large, but not nearly so large as  $\beta_n$ . Hence  $\sin \delta_n$  oscillates rapidly in the region  $0 < n < m$ , also, since  $\delta_{2n} = \beta_{2n}$ ,  $\delta_{2n+1} = \gamma_{2n+1}$ ,  $\sin \delta_{2n}$  and  $\sin \delta_{2n+1}$  will be independent functions. We can therefore replace  $\sin^2 \delta_n$  and  $\sin 2\delta_n$  in

expression (32) by their mean values of  $\frac{1}{2}$  and 0 respectively. We thus have\*

$$\begin{aligned} Q_D^1 &= 2\pi k^{-2} \sum_0^{\infty} \frac{1}{2} n \\ &= \frac{1}{2} \pi k^{-2} (m^2 + m) \end{aligned} \quad (34)$$

For values of  $n$  greater than  $m$ , the series (32) may be summed directly to give the value of  $Q_D^2$  as  $n$  increases, a point is reached where  $\beta_n \sim \gamma_n$  and the series begins to converge quite rapidly.

Values of  $Q_D$  are so obtained for different values of  $k$ , and are then used in formula (26) in order to take account of the Maxwellian distribution of velocities. Substitution in relations (25) and (24) then gives the value of the mobility at the required temperature. The final result of the calculation is that  $k = 12$  cm./sec per volt/cm at 760 mm Hg and 20° C. This is to be contrasted with the most recent value, 21.4, obtained by Tyndall and Powell†. This discrepancy might at first sight seem surprising, in view of the fact that Hassé‡ obtained nearly the experimental value by using classical theory and an assumed field based on the interaction of two helium atoms. He takes a single interaction energy

$$V_n = v_n - C/r^4, \quad (35)$$

where  $v_n$  is the interaction energy (1) between two helium atoms. Reference to figs 1 and 3 shows that this potential cannot be correct, and it has no justification in quantum theory, as the effect of nuclear symmetry extends the range of the potentials  $v_n$ ,  $v_s$  beyond that of  $v_n$ . Since a larger range interaction means a smaller mobility, it is to be expected that the mobility calculated above will be smaller than that calculated by Hassé. A check on the method given above for the calculation of the mobility was made by computing the mobility for the field  $V_n$  used by Hassé, the value 24 was then obtained, in fair agreement with the value 26 deduced by him from classical theory§. Moreover, the quantum theoretical calculation indicates an unvarying value of 24 for the mobility, provided that the fields  $v_n$  and  $v_s$  fall off sufficiently rapidly: thus if  $v_n$  and  $v_s$  have become small compared with the polarization

\* If the fields  $V_n$ ,  $V_s$  fall off rapidly at the point  $r_0$ , we have  $m \simeq kr_0$  and then equation (34) merely corresponds to the fact that at not too low velocities quantum and classical theory both give  $Q_D = \frac{1}{2} \pi r_0^2$  for a hard sphere of radius  $r_0$ .

† 'Proc. Roy. Soc., A', vol. 134, p. 125 (1931).

‡ 'Phil. Mag.', vol. 1, p. 139 (1926).

§ Hassé and Cook ('Phil. Mag.', vol. 12, p. 554 (1931)) have deduced values of 21 and 19.7 on the arbitrary assumption that the probability of charge transfer at any gas-kinetic collision is one-half, in an attempt to take into account exchange processes.

term  $C/r^4$  at a distance less than  $6a_0$ , whatever their value inside this point, the mobility is determined entirely by the polarization field beyond the point  $6a_0$ , and a value of 24 for the mobility is obtained.

However, it is important to obtain as much information as possible about the interaction in order to trace the discrepancy. It might well be that the formula (28), derived by a perturbation method, is inaccurate in the intermediate region where the two terms are comparable in magnitude. Another method was therefore used to calculate  $V_s$  and  $V_a$  which should give accurate results. The method is that used by Hylleraas\* for calculating exactly the interaction of a hydrogen atom and a proton. Reference to the papers of Pauling and Majorana shows that at the distances at which we are concerned the interaction energy terms  $V_s$  and  $V_a$  arise almost entirely from a single electron and the possibilities of nuclear interchange. In short, we may treat the interaction of He and He as a two-centre problem involving a single electron, but with an effective nuclear charge of  $Z$  instead of 1 where  $Z$  is taken as 1.5 (the value 1.45 leads to the correct polarization at large distances, while the value 1.48 is used in the wave function (29)). Now Hylleraas has shown that the interaction energy for different nuclear separations  $r$  is given by  $Z^3 f(r/Z)$ , so that we have merely to carry out Hylleraas' calculations for hydrogen for much greater nuclear separations, and then convert the values to those for helium by taking  $Z = 1.5$  in the expression  $Z^3 f(r/Z)$ . For  $V_s$  we have  $n = 1$ ,  $l = 0$ ,  $m = 0$ , and for  $V_a$   $n = 2$ ,  $l = 1$ ,  $m = 0$  in Hylleraas' calculations†. The computations must be carried out with high accuracy, as the energies  $V_s$ ,  $V_a$  are very small compared with the total energy. The resulting values obtained for  $\text{He}_2^+$  for various distances are given in Table IV in units of  $e^2/2a_0$ —the atomic unit of energy (13.53 volts), and in fig. 3 the corresponding values (denoted by a  $X$ ) for  $\text{He}_2^+$  are shown, together with the curves which were obtained with the use of the formula (28) based on the perturbation method, and which were used in obtaining the mobility.

It will be seen that the agreement is very satisfactory, and makes it improbable that any serious error can arise in the above calculations from inaccuracy in the form of  $V_s$  and  $V_a$ . It is difficult to see where the error lies, and it would be of great interest to obtain further experimental values over a considerable temperature range, if this should be possible. It would also be of great

\* 'Z. Physik,' vol. 71, p. 739 (1931).

† It may be useful here to mention that there are a few minor misprints in Hylleraas' paper. In particular, it is obvious that a term  $-2\sqrt{C}$  has been omitted in printing the first term of the infinite determinant (28).



value to measure the free path of helium ions in helium under gas-kinetic conditions. The cross-section effective in the collision of a helium ion with a helium atom was calculated for the above interaction by methods similar to those used in calculating the mobility cross-section. A value of  $10.3 \times \sqrt{2}a_0 = 14.6 a_0$  was obtained for the collision radius for a velocity corresponding to a temperature of  $20^\circ \text{C}$ .

It is interesting to note the effect of charge transfer in determining the values obtained above. The potentials  $V_a$  and  $V_s$  taken separately give values of 25 and 12 respectively for the mobility, as compared with the value 12 obtained if the fields  $V_a$  and  $V_s$  are combined, in the manner already

Table IV -- Interaction Energy of H and  $\text{H}^+$  for Different Nuclear Separations

$r/a_0$	$V_a$	$r/a_0$	$V_s$
3.3700	0.1886	2.9523	-0.1582
5.1682	0.0392	4.9810	-0.0405
7.0842	0.00706	7.0293	-0.0107
		8.0371	-0.00455
9.0533	0.00084	9.0430	-0.00169
11.0412	-0.00008	11.0386	-0.00059
13.0349	-0.00012	13.0344	-0.00020

shown, to take account of charge transfer. In the latter case, the mobility is mainly determined by the field  $V_s$ , which is larger in magnitude inside a distance  $8a_0$  than the field  $V_a$ , while the field beyond  $9a_0$  contributes comparatively little. For the collision cross-section, however, the effect of taking charge transfer into account is seen, on calculation, to be quite small.

### § 3 Calculation of Cross-Sections for Various Laws of Force

In considering the collision of helium atoms, the effect of the attractive van der Waals potential could be neglected except at quite low temperatures, but for all other gases and vapours the attractive field plays an important and usually predominant part—at least in the calculation of free paths. The asymptotic form of the attractive potential is known to be  $V \sim -C/r^6$ , and the constant  $C$  is usually so large that, for reasons we shall see, the repulsive field cannot affect the result. We shall now calculate the collision cross-section for a potential of the form  $-Cr^{-6}$ . The total cross-section  $Q$  is given by

$$Q = \frac{4\pi}{k^2} \sum_n (2n+1) \sin^2 \delta_n \quad (36)$$

and in all the cases of interest a large number of terms  $n$  are required in this series. However, the summation may be carried out in much the same way as the calculation of  $Q_D$  in § 2

The phases  $\delta_n$  are given by Jeffreys's approximation in the form\*

$$\delta_n = \int^{\infty} \left( k^2 - \frac{n(n+1)}{r^2} \right)^{\frac{1}{2}} dr - \int^{\infty} \left( k^2 - \alpha V - \frac{n(n+1)}{r^2} \right)^{\frac{1}{2}} dr, \quad (37)$$

where the lower limits of integration are the zeros of the respective integrands, and  $\alpha$  is written for  $8\pi^2 M \hbar^{-2}$ . For large  $n$  we have

$$\begin{aligned} \delta_n &\simeq \int \left( k^2 - \frac{n(n+1)}{r^2} \right)^{\frac{1}{2}} \left( 1 - 1 + \frac{\frac{1}{2}\alpha V}{k^2 - \frac{n(n+1)}{r^2}} \right) dr \\ &= \int \frac{\frac{1}{2}\alpha V}{\left( k^2 - \frac{n(n+1)}{r^2} \right)^{\frac{1}{2}}} dr, \end{aligned} \quad (38)$$

a result which is also given by Born's formula, as may be shown† by using the asymptotic expression for the Bessel function for  $n$  large. Jeffreys's approximation may therefore be used for all values of  $n$ .

Now if  $\alpha V = -C/r^2$ , we have for large  $n$ ,

$$\begin{aligned} \delta_n &\simeq \frac{1}{2}C \int_a^{\infty} \frac{1}{r^2 \left( k^2 - \frac{n(n+1)}{r^2} \right)^{\frac{1}{2}}} dr \\ &= \frac{1}{2} \frac{C}{k} \int_a^{\infty} \frac{1}{r^{2-1} (r^2 - a^2)^{\frac{1}{2}}} dr, \end{aligned}$$

where  $a = (n + \frac{1}{2})/k$ . On integration we easily obtain

$$\begin{aligned} \delta_n &= \frac{C}{2ka^{2-1}} f(s) \\ &= \frac{Ck^{2-2}}{2(n + \frac{1}{2})^{2-1}} f(s), \end{aligned} \quad (39)$$

\* The formula given here is slightly different in form to that given in paper I. The present form is much more convenient and accurate when used for the computation of phases.

† Massey and Smith, *loc. cit.*

where

$$f(s) = \frac{s-3}{s-2} \frac{s-5}{s-4} \quad \frac{1}{2} \frac{\pi}{2}, \quad s \text{ even},$$

$$\frac{s-3}{s-2} \frac{s-5}{s-4} \quad \frac{3}{2}, \quad s \text{ odd},$$

$$1, \quad s=3, \quad \frac{\pi}{2}, \quad s=2 \quad (40)$$

The use of more exact methods in a number of cases showed that this formula holds quite accurately if  $\delta_n$  is less than 0.5, and then  $\sin \delta_n \simeq \delta_n$ . Hence, if  $m$  is that value of  $n$  for which  $\delta_n = 0.5$ , we have for the contribution to the cross-section  $Q$  from  $n > m$

$$\begin{aligned} \frac{4\pi}{k^2} \sum_{n=m}^{\infty} (2n+1) \sin^2 \delta_n &= \frac{8\pi}{k^2} \sum_{n=m}^{\infty} \frac{C^2}{4} \frac{k^{2n-4}}{(n+\frac{1}{2})^{2n+3}} f^2 \\ &\simeq 2\pi C^2 k^{2m-6} f^2 \int_m^{\infty} (n+\frac{1}{2})^{-2n+3} dn \\ &= \frac{2\pi C^2 k^{2m-6}}{2s-4} (m+\frac{1}{2})^{-2s+6} f^2 \\ &= \frac{4\pi}{s-2} \frac{(m+\frac{1}{2})^2}{k^2} \delta_m^2, \text{ using (39),} \\ &= \frac{\pi}{s-2} \frac{(m+\frac{1}{2})^2}{k^2}, \end{aligned} \quad (41)$$

since by definition  $\delta_m = 0.5$ . Now the contribution to the total cross-section  $Q$  from  $n < m$  is given approximately—using the method of § 2—by

$$2\pi \frac{m^2}{k^2} \quad (42)$$

The total cross-section  $Q$  is therefore nearly equal to

$$\pi \left( 2 + \frac{1}{s-2} \right) \frac{(m+\frac{1}{2})^2}{k^2},$$

since  $m$  is large, and using (39) with the fact that  $\delta_m = 0.5$ , we obtain finally

$$Q = \pi \frac{2s-3}{s-2} f^{2/(s-1)} \left( \frac{C}{k} \right)^{2/(s-1)} \quad (43)$$

The accuracy of this formula was investigated more carefully by breaking the summation (36) into three parts instead of two. Particular values of  $s$ ,  $C$ , and  $k$  were chosen, and for each the phases were determined in the intermediate region by numerical integration using Jeffreys's formula (37), the

summation (36) then being carried out in this region by direct calculation. The contribution to  $Q$  from the region of quite large and quite small phases was determined as before. The results indicated that the formula (43) should give the total cross-section accurate to within about 5%.

The relation (43) shows that the collision cross-section is proportional to  $(C/v)^{2/(s-1)}$ , where  $v$  is the velocity of the colliding particles. Now classical theory predicts for the intensity of the scattering at any particular angle the same dependence on the velocity and the law of force\*. On classical theory, however, the total cross-section tends to larger and larger values as smaller and smaller deviations are taken into account in the measurements, whereas quantum theory gives the perfectly definite value (43).

Let us now apply the formula to the determination of the mean free path in argon at room temperature. The average kinetic energy of the atoms then corresponds to  $k = 10/a_0$ . Now the attractive van der Waals force between the atoms has the value  $31.8 \times 10^{-10} (r/a_0)^{-8}$  erg,† so that we have  $s = 6$ ,  $C = 5.5 \times 10^6/a_0^3$ . Substituting in (43) we obtain  $Q = 365 \pi a_0^2$ , and hence a collision radius of  $19.1 a_0$  for a pair of colliding argon atoms. This corresponds to the collision of two rigid spheres, each of diameter  $19.1 a_0/\sqrt{2} = 13.5 a_0$ .

Again, for water vapour at room temperature, the attractive force is effectively that between two dipoles, and from the dipole moment of the water molecule we deduce  $C = 960/a_0^3$ ,  $s = 3$ , while  $k = 6.7/a$ . Substitution in (43) gives  $Q = 4.40 \pi a_0^2$ . As for argon, this value has not been averaged over the Maxwellian distribution of velocities.

It is thus possible, with the use of formula (43), to interpret immediately experimental results obtained by molecular ray methods. It is important in making these measurements that the angular resolution of the apparatus be sufficient to give accurate values. The method of ensuring this was discussed in paper I, and although the considerations introduced there were illustrated by reference to the rigid sphere model, the same considerations apply to the above cases where  $r_0$  is given by  $r_0 = \sqrt{(Q/2\pi)}$ ‡. In the foregoing we have not taken into account the repulsive field, but its effect can be neglected if the zero of the interaction energy falls appreciably inside the point  $r_0$ , a condition satisfied for most atoms. Thus for argon  $r_0 = 13.5 a_0$ , whereas the zero falls at about  $7a_0$ .

\* Darwin, 'Phil. Mag.', vol. 27, p. 499 (1914).

† Lennard-Jones, 'Proc. Phys. Soc.', vol. 43, p. 461 (1931).

‡ It should be noted that in paper I, and in the present paper,  $r_0$ , for the collision of two rigid spheres, is the sum of the classical radii of the two spheres.

*Summary*

It has been shown in a previous paper that it is important to use the quantum theory of collisions for the determination of the laws of interaction between atoms at large distances from observations of transport phenomena and free paths. In the present paper, the form and magnitude of the interaction energy between two helium atoms is discussed in terms of observations of the viscosity of helium at different temperatures. The interaction between a helium atom and a helium ion is considered with reference to the mobility of helium ions in helium. Finally, formulæ are obtained which facilitate the determination of the law of force between atoms at large distances by direct measurements of free paths.

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*Studies on Gas-Solid Equilibria Part V -Pressure-concentration Equilibria between Silica Gel and (1) Oxygen, (2) Nitrogen, (3) Mixtures of Oxygen and Nitrogen, determined isothermally at 0° C.*

By BERTRAM LAMBERT and DAVID H. P. PEEL

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Experiments carried out on the adsorption of (1) pure oxygen and (2) pure nitrogen by a sample of silica gel, at 0° C., and at pressures up to 1 atmosphere, have shown that the adsorption is directly proportional to the pressure in each experiment.

The pressure-concentration isothermals are reproducible, completely reversible, and linear (except for a slight divergence at low pressures) throughout their whole length, there is therefore no sign of the silica gel approaching saturation with either oxygen or nitrogen at atmospheric pressure.

The study of the adsorption by the same silica gel from mixtures of oxygen and nitrogen (at 0° C. and within a total pressure range of 1 atmosphere) appeared, therefore, to offer a good opportunity of obtaining information, under simple conditions, on the problem of adsorption from binary gas mixtures.

The silica gel used was a sample from the same batch as that employed in the previous parts of these investigations.

A satisfactory study of adsorption from a binary gas mixture will involve measurements giving —

- (1) Accurate and reproducible pressure-concentration isothermals for each of the *pure gases* in contact with the adsorbent,
- (2) Accurate and reproducible isothermals for *mixtures*, showing the *partial pressures* of the components plotted against the amounts adsorbed at equilibrium

It is, of course, essential that the intrinsic adsorptive power of the adsorbent should undergo no variation during the whole course of the investigation

The determination of the partial pressures of a binary gas mixture in equilibrium with a solid adsorbent requires an accurate knowledge of the composition of the gaseous phase at equilibrium. A sample of the equilibrium gas mixture must be withdrawn from the adsorption system and, if this sample is to represent the true equilibrium gas phase, it must be withdrawn without disturbing the equilibrium at the surface of the solid. This gas sample must then be measured and analysed by a method which ensures a uniformly high degree of accuracy

The necessity for high accuracy in this operation is obvious when it is pointed out that errors at this stage are magnified many times in applying the results of the analysis to the determination of the true partial pressures in the equilibrium gas phase and to the amounts of each separate gas adsorbed at equilibrium

The difficulties of technique inherent in an investigation of this kind are probably responsible for the lack of reliable published data on the subject of adsorption from gas mixtures

The experimental difficulties have now been overcome completely and the application of a technique accurate at *all* stages has enabled reliable and repeatable results to be obtained for the adsorption from mixtures of oxygen and nitrogen by a sample of silica gel

The experimental procedure adopted in this investigation may be summarized as follows —

A known weight (*in vacuo*) of the activated silica gel was placed in an all-glass, sealed apparatus consisting of five parts, separated, where necessary, by stop-cocks —

- (A) *The calibrated gas-volume burette* for the accurate measurement of the gases to be introduced into or withdrawn from the adsorption system

- (B) *The distributor vessel* to enable the measured volumes of gases to be introduced into the adsorption system and also to enable samples of gas to be withdrawn from the adsorption system for subsequent measurement and analysis
- (C) *The adsorption system*—a vessel consisting of two compartments, one containing the solid adsorbent and the other (from which the sample of equilibrium gas mixture is withdrawn) separable from the first by means of a magnetically-operated internal valve, by this means it was possible to remove a sample of the true equilibrium gas phase without disturbing the gases adsorbed on the solid adsorbent
- (D) *The mercury manometer* for the measurement of equilibrium pressures This was attached to the adsorption system (gel compartment), the mercury being brought to a fixed point on this side of the manometer so that the volume of the adsorption system was invariable
- (E) *The gas analyser* used in conjunction with (A) for the analysis of the sample of equilibrium gas mixture after withdrawal from the adsorption system.

The adsorption system was maintained, throughout the whole period of each experiment, at 0° C by surrounding it with a large Dewar vessel (with drainage tube) filled with broken ice

The gas-volume burette and the manometer were surrounded by water jackets maintained at a temperature of 20° C ( $\pm 0.01^\circ$ ) Gas volumes could be read to 0.001 c.c. and pressures to 0.02 mm. of mercury by methods described below

The gas analyser was capable of giving uniformly reliable analyses of oxygen-nitrogen mixtures, the error being less than 0.1% in samples of 6 to 8 c.c.

The whole apparatus was thoroughly evacuated by means of a mercury vapour pump while the silica gel was heated for several hours at 150° C The residual pressure in the apparatus after this and all subsequent evacuations was less than  $10^{-4}$  mm. of mercury as measured by a McLeod gauge

After the regular procedure of "washing out" the apparatus,\* the volume (free gas space) of the adsorption system at 0° C was determined by the

\* A regular procedure was followed before the introduction of any gas or gas mixture into the adsorption system After thorough evacuation, during the last hour of which the silica gel was heated at 150° C, the cold system was "washed out" twice with the gas (or gases) to be used, the system being thoroughly evacuated and the silica gel heated for 1 hour at 150° C after each of these operations

introduction of measured volumes of pure helium, the pressure on the manometer being read after each addition, until the final pressure was about 1 atmosphere. The separate values thus obtained for the free gas space of the adsorption system did not vary more than 0.02%. Careful preliminary experiments carried out over a range of temperatures and pressures had shown that the adsorption of helium by silica gel at 0°C was negligible.

The true volume of the adsorption system being now known, after the usual preparation, measured volumes of oxygen and nitrogen were introduced and the (total) equilibrium pressure read off on the manometer. It was found, contrary to expectation, that the attainment of equilibrium between the silica gel and mixtures of oxygen and nitrogen was rapid and a pressure read within half an hour of the introduction of the gases into the adsorption system underwent no change over a period of several days. Nevertheless, 12 hours was allowed after the introduction of gas before the equilibrium pressure was read on the manometer.

The gel compartment of the adsorption system with the attached manometer was then shut off from the rest of the adsorption system (by closing the magnetically-operated internal valve) and a sample of the equilibrium gas phase withdrawn into the distributor vessel. This sample of the true equilibrium gas phase was now measured accurately in the gas-volume burette before transference to the analyser. In the analyser the measured sample of mixed gases was submitted to the action of red-hot pure iron (electroplated on a platinum spiral which could be electrically heated—details of the manipulation are given below). The oxygen having been completely removed by this process, the residual nitrogen was transferred back to the gas-volume burette for measurement.

These operations, all conducted within a sealed glass apparatus in which the gases could be transferred from one part to another by means of mercury, made it possible to get accurate and reliable values for—

- (1) the volume (free gas space) of the adsorption system,
- (2) the volumes of each separate gas added to the adsorption system,
- (3) the total pressure of the mixed gases in equilibrium with the gel at 0°C,
- (4) the composition of the gaseous phase at equilibrium.

It was thus possible to calculate the exact amounts of each gas adsorbed from a mixture of oxygen and nitrogen and the exact partial pressures of each of the gases in equilibrium with the gel under these conditions.



Experiments were conducted within the total pressure limits of 1 atmosphere and the results showed a high degree of repeatability. Accurate and reproducible isothermals could be drawn showing values of the equilibrium partial pressures of oxygen and nitrogen plotted against the amounts of each separate gas adsorbed. A comparison of these isothermals with those for pure oxygen and pure nitrogen shows how the adsorption of each gas is affected by the presence of the other.

The apparatus used in this work is shown diagrammatically in fig 1. For the purpose of this description it is conveniently divided into five sections as in the introduction above, the five sections are designated by the letters A, B, C, D, E, both in the diagram and in the description.

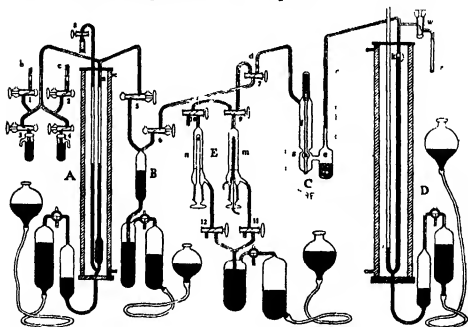


FIG 1

#### (A) *The Calibrated Gas-volume Burette*

This was of the type generally used when gas volumes are measured at atmospheric pressure, it consisted of a measuring limb and a balancing limb connected together and attached, at the bottom end, to a mercury reservoir. The graduated part of the measuring limb had the same internal diameter (5 mm) as the balancing limb, meniscus corrections were thus avoided and

gas contained in the measuring limb was at atmospheric pressure when the mercury levels in the two limbs were brought to the same height, with the balancing limb open to the air

The burette was mounted vertically in a jacket fitted with plate-glass windows back and front. It was maintained at a temperature of  $20^{\circ}\text{C}$  ( $\pm 0.01^{\circ}$ ) by the rapid circulation through the jacket of water from a large constant-temperature bath, the thermostatic control of the bath being adjusted from time to time to compensate for alterations in the room temperature

The capacity of the measuring limb of the burette was 12 c.c. and it was graduated every 0.02 c.c. It was very carefully calibrated at  $20^{\circ}\text{C}$  by the weighed mercury method

The burette was read by means of a reading microscope fitted with a micrometer scale in the eye-piece. The magnification of the microscope was adjusted so that 20 micrometer scale divisions exactly covered the space between the centres of two adjacent graduation marks when the burette was viewed through the microscope, volume readings could therefore be made to 0.001 c.c., or less, by estimation. The reading microscope was mounted on a carriage which moved vertically, by rack and pinion action, on an accurately-machined steel bar fixed to the front of the jacket, the microscope carriage was also provided with a true horizontal movement which was brought into action after locking the carriage in a given vertical position

This precision reading device attached to the burette jacket was found to be much superior to a cathetometer, there was a complete freedom from all detectable errors in reading the position of the mercury meniscus on the etched burette scale and gas volume readings could be repeated with certainty to less than 0.001 c.c. Reflected light effects were removed from each mercury meniscus by a black ebonite shroud surrounding the glass tube and brought to a position about 1 mm. above the mercury level, this enabled the mercury meniscus to be viewed as a black silhouette when illuminated by a suitably diffused light placed behind the water jacket

The mercury reservoir was connected to the burette by glass tubing and its position was thus fixed. The alterations of pressure necessary to work the reservoir were obtained by compressing or rarefying the air in contact with the mercury by means of an external mercury compressor as shown in the diagram. The movable compressor-reservoir was carried in a cradle moving in a long vertical slide and a fine adjustment was provided so that the levelling of the mercury in the burette limbs could be carried out satisfactorily to within 0.02 mm. The T-stop-cock between the burette reservoir and the

mercury compressor allowed of variations in the amount of air in the intervening air-buffer and so permitted the application of a wide range of positive and negative pressures to the mercury in the burette reservoir. This method of working a mercury reservoir was found to be very satisfactory and was used on all those employed throughout the apparatus.

The upper ends of the burette limbs were made of capillary tubing of 1 mm. bore, and the zero mark of the burette was an etched line *a* on the capillary part of the measuring limb. The balancing limb terminated in a stop-cock 8 opening to the air through a protecting tube containing soda-lime and active charcoal granules. Above the zero mark the capillary tubing branched at a Y-junction, one side being connected with the distributor vessel B (through the stop-cock 5) and the other with the oxygen and nitrogen storage vessels (through stop-cocks 1 and 2), the gas storage vessels were connected by capillary tubing, the oxygen storage vessel being sealed on at *b* and the nitrogen storage vessel at *c*. Stop-cocks 3 and 4 communicated with the open air through mercury cups with side-tubes as shown. All connecting tubes were made of 1 mm. bore tubing.

With suitable pressure adjustments on the mercury reservoirs of the burette and the distributor vessel, the system of stop-cocks allowed the complete removal of all gas and its replacement by mercury in the capillary connections between these vessels and the gas storage vessels, stop-cocks 1 to 5 being then closed, reduction of the pressure on the burette reservoir, followed by the opening of stop-cock 1 (or 2), allowed gas to be drawn from the oxygen (or nitrogen) storage vessel into the burette. Finally, the opening of stop-cock 3 (or 4) and the application of pressure (by means of a hand-bellows) to the mercury in the attached mercury cup, allowed the oxygen (or nitrogen) left behind in the connecting tube to be pushed into the burette (and above stop-cock 1 (or 2)) and the mercury level to be adjusted at the zero mark *a* on the burette.

A suitable volume of pure oxygen or nitrogen could thus be transferred from its storage vessel to the burette without introducing any impurity or leaving any traces behind in the connecting tubes. The mercury levels could then be adjusted so as to bring the gas to atmospheric pressure and its volume accurately measured. All measured volumes of gases were reduced to standard temperature and pressure.

#### *(B) The Distributor Vessel*

This consisted of a cylindrical glass bulb (of about 30 c.c. capacity) provided, at its lower end, with a mercury reservoir to which a wide range of positive

and negative pressures could be applied by the method described under A, the upper end terminated in a Y-junction, of capillary bore, connecting it on one side with the gas-volume burette A (through stop-cock 5) and on the other side with either the adsorption system C or the gas analyser E (through stop-cock 6 and the appropriate limb of the three-way stop-cock 7). All the connecting tubes were made from capillary tubing of 1 mm. bore and all the stop-cocks had accurately-fitting stoppers with the same bore.

By suitable adjustment of pressures on the mercury reservoirs concerned, it was possible to transfer gas completely—following it with mercury and avoiding the trapping of bubbles of gas—(1) from the distributor vessel B to the gas-volume burette and *vice versa*, (2) from the distributor vessel B to the gas analyser E and *vice versa*, (3) from the distributor vessel B to the adsorption system C. In this last operation the mercury following the gas was stopped at the fixed mark *d* which was one of the two limits defining the volume of the adsorption system.

The distributor vessel was also used to remove gas from the adsorption system, after the gas thus removed had been transferred to the burette for measurement, the gas left in the tubes and stop-cocks between the distributor vessel and the adsorption system was pushed back into the latter section and the mercury again brought to the fixed mark *d*.

### (C) *The Adsorption System.*

This section of the apparatus consisted of two compartments—referred to, throughout this description, as the gel compartment and the gas compartment. The gel compartment *e* contained the silica gel which was in the form of small granules, it was attached to the manometer D by capillary tubing and to the gas compartment *f* by tubing of wide (1 cm.) bore. The upper end of the gas compartment was connected, by capillary tubing, with the distributor vessel B (through the stop-cocks 7 and 6).

A magnetically operated internal valve which served, when required, to close the connection between the two compartments was made from light glass tubing blown into a bulb at *h* and widened at the top so as to enclose a hollow soft-iron cylinder. It was sealed at both ends and rested loosely in the gas compartment which was shaped at its upper end so as to hold the valve in an approximately vertical position.

When the valve was resting in its normal position, as it is shown in the diagram, the gel and gas compartments were open to one another and there

was no interference with the easy diffusion of gas throughout both compartments. The application of an electromagnet at the top of the gas compartment caused the valve to rise and the bulb *h* to be held in close contact with a ground seating *g* on a constricted part of the gas compartment. With the valve in this position, the connection between the two compartments was closed. The valve readily fell away from the ground seating when the electromagnet was switched off and so restored the open connection between the two compartments. When the valve was closed, gas could be withdrawn from the top section of the gas compartment into the distributor vessel B without appreciably affecting the pressure indicated by the manometer in the gel compartment.

In the experiments with gas mixtures, the use of this device ensured that a sample of gas (for the determination of the partial pressures of the two constituents) could be withdrawn without disturbing the equilibrium at the surface of the silica gel during the actual withdrawal operation and that the sample withdrawn was really representative of the true equilibrium gas phase.

The adsorption system was maintained at 0° C, throughout each experiment, by surrounding it with a large Dewar vessel filled with broken ice, the Dewar vessel was constructed with a drain-tube through the bottom.

The volume of the adsorption system was the gas space between the two fixed points *d* and *k* each defined by a mercury meniscus. A "dead-space" correction was made, throughout the work, for the very small fraction of the volume which was outside the ice-bath.

#### (D) *The Mercury Manometer*

This consisted of a long U-tube of internal diameter 8 mm. throughout, one limb being connected to the adsorption system by a short length of capillary tubing, the other limb being open to the air through a protecting tube containing soda-lime and active charcoal granules. The bottom of the U-tube was connected to a mercury reservoir to which a wide range of positive and negative pressures could be applied by the method described under A.

A pressure reading on this manometer involved three operations: (1) adjusting the pressure on the mercury reservoir so as to bring the mercury level to the fixed position *k* in the limb connected to the adsorption system; (2) measuring the exact difference in height between the mercury levels in the two limbs, (3) reading the standard barometer. The necessary temperature corrections were made so as to bring all pressure measurements to values expressed in millimetres of mercury at 0° C.

The manometer was mounted in a jacket made with machined sides and base of duralumin and having plate-glass windows back and front. Inside the jacket, and immediately behind the manometer, there was mounted a thick plate-glass scale. This scale was made by the Société Genevoise and guaranteed to be accurate (to within 5 microns) at 20° C over the whole length of 800 mm. It was graduated in millimetres and the graduation marks were of an even thickness of 0.05 mm. The manometer and scale were maintained at a temperature of 20° C ( $\pm 0.01^\circ$ ) by the method described under A.

A high-precision reading device was mounted on the front of the manometer jacket, this was similar to that fitted on the gas-volume burette except that two reading microscopes were used. The magnification of the microscopes was adjusted so that 25 micrometer (eye-piece) scale divisions exactly covered the space between the centres of two adjacent millimetre graduations, when the graduated scale was viewed through the microscopes. Direct readings could therefore be made to 0.04 mm., or less, by estimation.

In setting up this section of the apparatus, great care was exercised to ensure that all the essential parts were quite rigid and vertical.

When each mercury meniscus was properly shrouded and suitable illumination provided behind the manometer jacket (*vide supra*), it was possible—using the vertical and horizontal movements of the microscope carriages—to read very accurately the position of each mercury meniscus on the graduated glass scale. For a given pressure in the adsorption system, the vertical distance between the two mercury levels could be measured with certainty to less than 0.02 mm. No meniscus correction was necessary as each mercury meniscus was in tubing of the same bore.

Accurate *barometer* readings were necessary for the calculation of equilibrium pressures and also of gas volumes. These were made on a standard Fortan barometer and the usual temperature corrections made to reduce the readings to 0° C. Independent readings were made on a U-tube barometer set up and read like the manometer described above. This barometer was made from Pyrex glass, thoroughly “baked out” *in vacuo* and filled by distilling pure mercury into it. The readings of the two barometers, corrected to 0° C, agreed to within 0.02 mm.

#### (E) *The Gas Analyser*

This was used in conjunction with the gas-volume burette for the accurate analysis of the samples of equilibrium oxygen-nitrogen mixtures.

After a sample of equilibrium gas mixture had been withdrawn from the adsorption system and transferred to the burette, the gas remaining in the connecting tubes and stop-cocks between the distributor vessel and the adsorption system was pushed back into the latter, the mercury level adjusted again at the fixed mark *d* and the stop-cock 7 closed. The measured volume of oxygen-nitrogen mixture was then transferred, via the distributor vessel, to the gas analyser E.

In one part, *m*, of the gas analyser, the oxygen-nitrogen mixture was submitted to the action of red-hot, pure iron (electroplated on a spiral of platinum wire which could be heated electrically). This resulted in the conversion of the oxygen into iron oxide and its complete removal from the gaseous phase. Small traces of carbon dioxide and water vapour were necessarily produced in this operation since the oxygen-nitrogen mixture contained grease vapour owing to its passage through grease-lubricated stop-cocks, these impurities were removed from the residual nitrogen in another part, *n*, of the gas analyser, where it was left for 12 hours in contact with a Pyrex glass rod *o* coated with a thin layer of fused potash. The pure nitrogen was finally transferred back to the burette, its volume determined and the composition of the original mixture calculated.

The accuracy of this technique was thoroughly tested by analysing oxygen-nitrogen mixtures of known composition—made from measured volumes of the pure gases withdrawn from the storage vessels. The difference between the known composition of a mixture and that determined as above, was always less than 0.1% even when the total volume of gas analysed was only 6 to 8 c.c.

The detailed construction of the gas analyser is clearly shown in the diagram. The three-way stop-cocks 9, 10, 11, 12, arranged as shown, and the large mercury reservoir (to which a wide range of positive and negative pressures could be applied) made it possible to transfer the whole of the gas to where it was required for the various operations, and to follow it by mercury so avoiding the trapping of gas bubbles in the connecting tubes and stop-cocks.

The pressure arrangements attached to the mercury reservoir also made it possible to produce a Torricellian vacuum in either of the vessels *m* or *n*, so that dissolved air could be completely removed from the mercury and adsorbed gas from the iron-coated platinum spiral. Since this spiral had to be reduced—by heating it in a stream of pure, dry hydrogen and then cooling it in the gas—before each analysis, it had to be heated to bright redness several times, *in vacuo*, in order to remove the adsorbed hydrogen. The dissolved or adsorbed

gas thus removed was pushed out by mercury through the open limb of stop-cock 10

The arrangement of the stop-cocks 9, 10, 11, 12, made it possible—by the use of the appropriate open limbs of the last three—(1) to reduce the iron-coated platinum spiral by heating it in a current of pure, dry hydrogen and cooling it in the gas stream, (2) to renew the potash on the Pyrex rod *c* by removing it at the ground junction, dipping it in molten potash and cooling it, after replacement, in a stream of pure dry nitrogen. After such operations and the subsequent removal of dissolved and adsorbed gases, as described above, the whole system could be filled with mercury and left ready to receive a measured volume of gas for analysis.

Each stop-cock used in the construction of the apparatus was carefully chosen so that mercury could be passed through it without trapping small gas bubbles. Each stopper had a bore of the same diameter (about 1 mm.) as the capillary tubes sealed on to the stop-cock barrels and the openings of these tubes coincided exactly with the ends of the capillary passage through the stopper. The stop-cocks were lubricated with Ramsay stop-cock grease (previously boiled *in vacuo*) and, after the stoppers had been turned many times in both directions, the excess grease accumulated in the bore was washed out with carbon tetrachloride, air was then drawn through the bore for several hours.

Great care was taken to ensure that the gases used in these experiments were of the highest possible purity. Before filling with gas, each storage vessel was thoroughly evacuated, heated and "washed out" several times with the gas to be stored in it. Each gas storage vessel contained a supply of pure phosphorus pentoxide with which the gas was left in contact for several weeks before use.

The nitrogen was made by heating pure sodium azide (purified by several recrystallizations) in a wide tube directly sealed to the storage vessel. No purification of this gas was necessary.

The oxygen was made by the electrolysis of a saturated solution of barium hydroxide (purified by many recrystallizations). The electrolytic cell was similar in design to that used by Lambert,\* but air was kept out of contact with the electrolyte in the cell reservoir by means of a buffer of oxygen. Since traces of hydrogen must be contained in electrolytic oxygen, owing to the solubility of this gas in the electrolyte, the oxygen was purified by circulating it over a red-hot platinum spiral and pure phosphorus pentoxide for 20 hours.

\* 'Trans. Chem. Soc.' vol. 101, p. 2058 (1912)



The stop-cocks of the purifying and storage vessel were lubricated with glacial phosphoric acid. Gas drawn from this vessel was passed through a narrow U-tube immersed in liquid air to remove traces of water vapour taken up by the oxygen in its passage through the stop-cock lubricated with phosphoric acid.

The helium used for the measurement of the free gas space of the adsorption system was made from thoriated and purified by circulating it for many hours over large quantities of active charcoal cooled to liquid air temperature. The 100% helium supplied, in sealed glass tubes, by the British Oxygen Company, was also used and found to be quite satisfactory for the purpose.

The pressure-concentration equilibria between the silica gel and (1) pure nitrogen, and (2) pure oxygen were first determined. Two independent experiments were carried out with each gas, the adsorption system being evacuated and the silica gel heated for 1 hour at 150° C between each experiment.

The results are given in Tables I and II. The weight (*in vacuo*) of the silica gel used in the experiments was 9.8812 gm. Amounts of gas adsorbed by this weight of gel are given in cubic centimetres at standard temperature and pressure, equilibrium pressures are expressed in millimetres of mercury at 0° C.

These results are plotted in fig. 2.

The results show quite clearly that the equilibria between (1) silica gel and nitrogen, (2) silica gel and oxygen are completely reversible and reproducible over the whole pressure range and that the adsorption, in each experiment, is proportional to the pressure except over the lower range of pressures. It is noticeable that the amount of nitrogen adsorbed is over 10% more than the amount of oxygen adsorbed at the same pressure.

#### *Adsorption from Oxygen-Nitrogen Mixtures*

Two separate and independent experiments were carried out on the adsorption shown by the silica gel when exposed to mixtures of oxygen and nitrogen, the adsorption system being thoroughly evacuated, and the silica gel heated for 1 hour at 150° C, between the two experiments.

Mixtures containing approximately equal quantities of the two gases were used throughout and the partial pressures of the two gases in the adsorption system were never far apart. The gases were mixed in the adsorption system—the oxygen being introduced before the nitrogen—in the determination of each “ascending” equilibrium point.

Table I—Silica Gel—Nitrogen Equilibria at 0° C

No	Volume of nitrogen adsorbed.	Equilibrium pressures	Volume adsorbed Pressure
1	1 937	78 15	0 025
2	3 738	151 73	0 025
3	5 798	238 80	0 024
4	7 921	327 39	0 024
5	10 123	421 42	0 024
6	12 318	517 25	0 024
7	14 614	617 55	0 024
8	16 858	716 42	0 024
9	16 050	680 72	0 024
10	13 427	564 63	0 024
11	11 117	463 97	0 024
12	8 777	363 39	0 024
13	6 402	262 96	0 024
14	4 201	170 73	0 025
15	2 127	85 58	0 025
16	0 457	18 09	0 025
17	2 480	100 42	0 025
18	5 249	214 93	0 024
19	7 728	318 85	0 024
20	10 322	430 11	0 024
21	12 922	542 40	0 024
22	15 292	647 02	0 024
23	16 754	712 21	0 024
24	17 911	763 49	0 024
25	15 324	648 53	0 024
26	12 438	521 17	0 024
27	9 462	392 62	0 024
28	6 840	281 83	0 024
29	4 228	171 78	0 025
30	2 886	116 63	0 025
31	1 311	52 69	0 025

The equilibrium points are numbered in the order in which they were obtained

First experiment { "Ascending" points—Nos. 1 to 8  
 "Descending" points—Nos. 9 to 16  
 Second experiment { "Ascending" points—Nos. 17 to 24  
 "Descending" points—Nos. 25 to 31

The adsorption system was completely evacuated between the first and second experiments.

About 12 hours were allowed for the establishment of equilibrium after the addition (or withdrawal) of gas, although careful experiments had shown that the equilibrium (total) pressure was reached in half an hour and showed no subsequent alteration over a period of several days.

The results of the two experiments are given in Tables III and IV. The equilibrium points are numbered in the order in which they were obtained, each oxygen point in Table III being obtained in the same experiment as the nitrogen point designated by the same number in Table IV. The amounts of gas adsorbed by the gel (9 8812 gm) are given in cubic centimetres at standard temperature and pressure, and the equilibrium partial pressures are given in millimetres of mercury at 0° C. The results are given in two tables so as

Table II—Silica Gel—Oxygen Equilibria at 0° C

No	Volume of oxygen adsorbed	Equilibrium pressures.	Volume adsorbed Pressure
1	1 512	68 65	0 023
2	3 206	147 78	0 023
3	5 065	234 97	0 023
4	6 914	323 30	0 021
5	8 925	420 29	0 021
6	10 966	519 33	0 021
7	12 486	592 95	0 021
8	14 042	669 65	0 021
9	15 239	724 87	0 021
10	16 331	783 00	0 021
11	14 543	694 16	0 021
12	13 341	635 49	0 021
13	12 109	574 17	0 021
14	10 504	495 88	0 021
15	8 890	417 97	0 021
16	7 281	339 80	0 021
17	5 536	257 15	0 023
18	3 703	189 71	0 023
19	2 296	104 20	0 023
20	1 386	61 97	0 023
21	0 440	18 18	0 024
22	2 269	105 27	0 023
23	4 633	215 61	0 021
24	6 823	319 16	0 021
25	8 900	420 24	0 021
26	10 981	520 15	0 021
27	12 659	602 24	0 021
28	14 602	698 02	0 021
29	14 026	669 58	0 021
30	11 321	537 51	0 021
31	8 778	413 46	0 021
32	6 427	301 20	0 021
33	4 298	199 30	0 022
34	2 098	96 56	0 023
35	0 665	29 51	0 023

The equilibrium points are numbered in the order in which they were obtained

First experiment { "Ascending" points—Nos 1 to 10  
 "Descending" points—Nos. 11 to 21  
 Second experiment { "Ascending" points—Nos 22 to 32.  
 "Descending" points—Nos 29 to 35

The adsorption system was completely evacuated between the first and second experiments

to show clearly how the adsorption of each gas is affected by the presence of the other

In Table V the results are arranged so as to show how the added amounts of the oxygen and nitrogen adsorbed at the twelve equilibrium points compare with the sum of the amounts of these gases when adsorbed singly at the same pressures. It is clear that the total amount of gas adsorbed by the silica gel from mixtures of oxygen and nitrogen is slightly less than the sum of the separate amounts of the pure gases adsorbed singly at the same pressures.

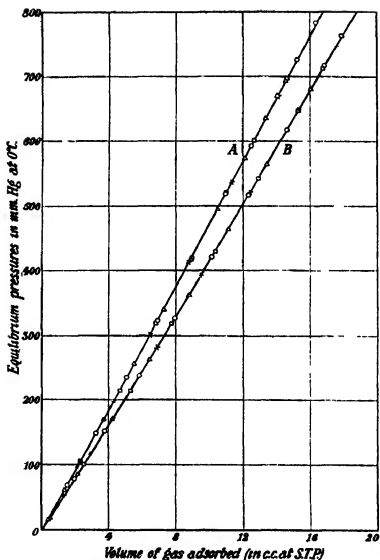


FIG. 2—Curve A oxygen, curve B nitrogen.

First experiments with each gas { "Ascending" points—○  
 "Descending" points—△  
 Second experiments with each gas { "Ascending" points—□  
 "Descending" points—×

The results are plotted in figs. 3 and 4, along with the isothermals for the pure gases, so as to show how the adsorption of each gas is affected by the presence of the other

Table III—Silica Gel—Oxygen Equilibria (in presence of Nitrogen)

No	Partial pressure of oxygen	Volume of oxygen adsorbed (mixture).	Volume of oxygen absorbed from pure oxygen at same pressure	Change in adsorption of oxygen due to the presence of nitrogen
1	89 45	1 345	1 336	+0 019
2	118 03	2 789	2 585	+0 204
3	170 48	4 109	3 709	+0 400
4	233 81	5 258	4 835	+0 423
5	107 63	2 787	2 360	+0 427
6	209 12	5 095	4 590	+0 575
7	286 60	6 930	6 148	+0 772
8	353 06	8 443	7 522	+0 921
9	330 73	7 939	7 075	+0 864
10	308 81	7 380	6 607	+0 773
11	287 89	6 916	6 170	+0 746
12	265 77	6 420	5 715	+0 705

Table IV—Silica Gel—Nitrogen Equilibria (in presence of Oxygen.)

No	Partial pressure of nitrogen.	Volume of nitrogen adsorbed (mixture).	Volume of nitrogen absorbed from pure nitrogen at same pressure.	Change in adsorption of nitrogen due to presence of oxygen.
1	59 50	1 346	1 485	-0 139
2	118 85	2 603	2 936	-0 334
3	172 39	3 701	4 238	-0 537
4	236 24	4 885	5 533	-0 648
5	112 36	2 339	2 780	-0 441
6	214 17	4 583	5 243	-0 660
7	296 32	6 339	7 300	-0 961
8	365 01	7 831	8 837	-1 006
9	343 00	7 293	8 285	-0 992
10	318 83	6 825	7 739	-0 914
11	297 10	6 372	7 219	-0 847
12	274 20	5 879	6 679	-0 800

First experiment      "Ascending" points—Nos. 1 to 4  
 Second experiment    { "Ascending" points—Nos. 5 to 8.  
                               "Descending" points—Nos. 9 to 12.

After the above experiments had been completed the adsorptive power of the gel for each of the pure gases was re-determined. It was found that the intrinsic adsorptive power of the gel for pure oxygen and pure nitrogen had undergone no change whatever, equilibrium points were obtained which fell exactly on the originally-determined isothermals.

The accuracy of the technique employed in this work and the reproducibility of the results in separate, independent experiments, seem to leave no doubt that, with the sample of silica gel used, the adsorption of oxygen from mixtures

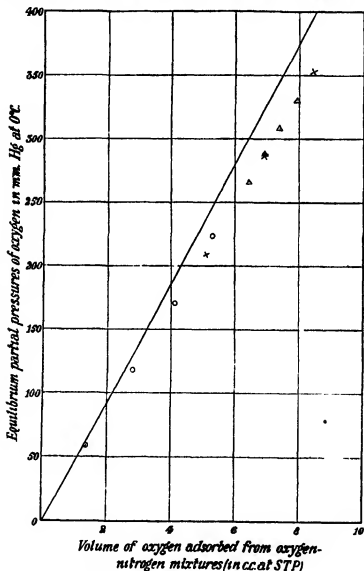


FIG. 3.—Pure oxygen isothermal.

of oxygen and nitrogen (at approximately equal pressures) is markedly higher than it is from pure oxygen. The adsorption of nitrogen is, on the other hand, definitely lower than it is from pure nitrogen. Although the silica gel showed

a greater adsorption from pure nitrogen than from pure oxygen (at corresponding pressures) this order is reversed when the gel is placed in contact with mixtures of approximately equal volumes of the two gases.

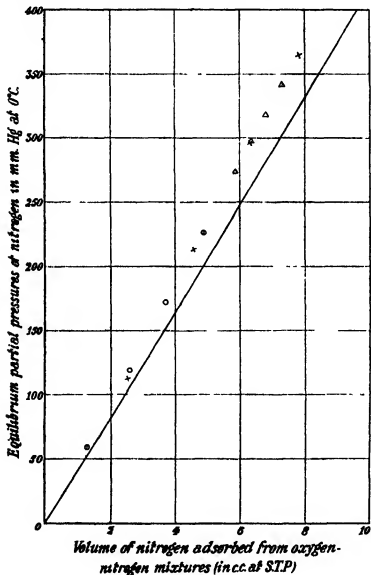


FIG. 4.—Pure nitrogen isothermal.

A discussion of these results is postponed until data are obtained for the adsorption by the same sample of silica gel from other binary gas mixtures.

Table V—Silica Gel—Oxygen-Nitrogen Equilibria at 0° C

No	Oxygen		Nitrogen		Totals.		Change on mixing
	Volume O <sub>2</sub> adsorbed, mixture	Volume O <sub>2</sub> adsorbed, single gas	Volume N <sub>2</sub> adsorbed, mixture	Volume N <sub>2</sub> adsorbed, single gas.	Total gas adsorbed, mixture	Total gas adsorbed, singly	
1	1 345	1 326	1 346	1 485	2 691	2 811	-0 110
2	2 789	2 585	2 602	2 936	5 391	5 531	-0 130
3	4 109	3 709	3 701	4 238	7 810	7 947	-0 137
4	5 288	4 835	4 885	5 533	10 173	10 368	-0 195
5	2 787	2 360	2 339	2 780	5 126	5 140	-0 014
6	5 085	4 520	4 583	5 243	9 678	9 763	-0 085
7	6 920	6 148	6 339	7 200	13 259	13 348	-0 089
8	8 443	7 522	7 831	8 837	16 274	16 359	-0 085
9	7 939	7 075	7 293	8 285	15 232	15 359	-0 128
10	7 380	6 607	6 825	7 739	14 205	14 346	-0 141
11	6 916	6 170	6 372	7 219	13 288	13 389	-0 101
12	6 420	5 715	5 879	6 679	12 299	12 394	-0 095

First experiment "Ascending" points—Nos. 1 to 4

Second experiment { "Ascending" points—Nos. 5 to 8  
"Descending" points—Nos. 9 to 12

Thanks are tendered to Imperial Chemical Industries for a grant which helped to defray the cost of this investigation

### Summary

(1) An accurate technique is described for the study of adsorption by solid adsorbents from binary gas mixtures.

This technique has been applied to the investigation of the adsorption by silica gel at 0° C. from mixtures of oxygen and nitrogen when these gases are present in approximately equal quantities

(2) An investigation of the adsorption of the *separate* gases at 0° C. has shown that, in each experiment, the adsorption is directly proportional to the pressure, except for a very slight divergence at pressures below 100 mm. The pressure-concentration isothermals show a complete reversibility over the whole range of pressures studied (up to 1 atmosphere), and they are reproducible throughout in separate independent experiments.

The adsorptive power of the silica gel is greater, at corresponding pressures, for pure nitrogen than for pure oxygen

(3) A study of the adsorption at 0° C. from *mixtures* of approximately equal quantities of these pure gases has yielded remarkable results—



- (a) The adsorptive power of the silica gel for oxygen is, in the presence of nitrogen actually greater (up to 15%) than when oxygen alone is present
- (b) On the other hand, the adsorptive power of the gel for nitrogen is less in the presence of oxygen than when nitrogen alone is present
- (c) The total amount of gas adsorbed from the mixture is always slightly less than the sum of the separate amounts of the pure gases absorbed singly at the same pressures

### *The Theory of Alloys in the $\gamma$ -Phase*

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#### 1. Introduction

Binary alloys in the  $\gamma$ -phase according to Endo† and Bernal‡ have especially large diamagnetic susceptibilities. It appears also that the Hall coefficients§ may be much greater for the alloys in this phase than for either of the constituent pure metals. It will be shown that it is possible to relate these properties to the crystal structure of the alloys, and to the fact that the composition within the  $\gamma$ -phase follows the Hume-Rothery|| electronic rule. For alloys in this phase the rule states that there are 21 loosely bound, or valency electrons, to every 13 atoms. A reason will also be shown to exist for this rather odd ratio of the number of approximately free electrons to the number of atoms.

The remainder of this section gives a summary of results already known which we shall require.

The wave-function of an electron moving in the periodic field of a lattice is of the form¶

$$\psi_k = e^{\frac{2\pi i}{a}(\mathbf{k} \cdot \mathbf{r})} u_k(\mathbf{r}), \quad (1)$$

† 'Sol. Rep. Imp. Univ. Tokyo,' vol. 14, p. 479 (1925).

‡ 'Trans. Faraday Soc.,' vol. 25, p. 367 (1929).

§ Richards and Evans, 'Phil. Mag.,' vol. 13, p. 201 (1932).

|| 'The Metallic State,' p. 328.

¶ Bloch, 'Z. Physik,' vol. 52, p. 555 (1929).

where  $u_{\mathbf{k}}(\mathbf{r})$  is periodic with the period of the potential field, and the three components of  $\mathbf{k}$  are the numbers used to specify a state. The  $\gamma$ -structures are based on a cubic lattice, so that it will be sufficient only to consider crystals with cubic symmetry.  $a$  is then the length of the side of a unit cell.

It is convenient to regard  $k_x, k_y, k_z$  as plotted along three rectangular axes, so that every state of an electron in the lattice is represented by a point in  $k$  space. The energy of a state associated with a point in  $k$  space may be determined to a first approximation by perturbing free electrons with the lattice field. For free electrons the energy is

$$E_{\mathbf{k}}^{(0)} = \frac{\hbar^2}{2ma^2} |\mathbf{k}|^2 \quad (2)$$

Since the momentum of a free electron is  $\hbar\mathbf{k}/a$ , and since owing to spin there are two states per  $h^3$  of phase space, it follows that the number of states per

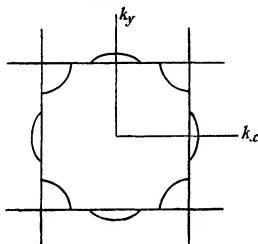


FIG. 1

unit volume of  $k$  space is  $2N/8$  where  $N$  is the total number of atoms in the crystal and 8 the number per unit cell. Hence the density of states per atom in  $k$  space is  $2/8$ .

It is a well-known result of the theory that the effect of the lattice field is to introduce planes in  $k$  space across which the energy changes discontinuously †. For example, for a two dimensional cubic lattice the energy is discontinuous across the sides of a square as shown in fig. 1. The curved line is a line of constant energy, such a line will represent the boundary between states which

† Brillouin, "Quantenstatistik," chap. 8.

are occupied and states which are unoccupied. Such a boundary is sharply defined at low temperatures. In a three dimensional lattice it forms a surface in  $k$  space which for convenience is usually referred to as the surface of the Fermi distribution.

The equations to the planes across which the energy may change discontinuously are given by the conditions

$$E_{\mathbf{k}+\mathbf{n}}^{(0)} - E_{\mathbf{k}}^{(0)} = 0, \quad (3)$$

where  $\mathbf{n}$  stands for the three integers ( $n_1, n_2, n_3$ ) each of which may be positive, negative, or zero. These planes, viz.,

$$n_1 k_x + n_2 k_y + n_3 k_z + \frac{1}{2} (n_1^2 + n_2^2 + n_3^2) = 0 \quad (4)$$

be parallel in  $k$  space to the set of parallel planes in the crystal associated with the Miller indices ( $n_1, n_2, n_3$ ).

If the potential of the lattice field be expressed in the form

$$V = \sum_{\mathbf{n}} V_{\mathbf{n}} e^{\frac{2\pi i}{a}(\mathbf{n} \cdot \mathbf{r})} \quad (5)$$

the discontinuity in the energy on crossing the plane ( $n_1, n_2, n_3$ ) in  $k$  space is given to the first approximation by

$$\Delta E_{\mathbf{n}} = 2e|V_{\mathbf{n}}| \quad (6)$$

This approximation to the energy gap becomes more accurate as the energy of the state corresponding to the point ( $\frac{1}{2}n_1, \frac{1}{2}n_2, \frac{1}{2}n_3$ ) in  $k$  space increases, i.e., as  $\frac{1}{2}n$  increases, because the greater the energy the more nearly do the wavefunctions of the states approximate the plane waves.

## 2 The Relative Magnitudes of the Fourier Coefficients of the Potential

As any crystal with cubic symmetry can be built up of interpenetrating simple cubic lattices, the potential may be written

$$V = \sum_{\mathbf{n}} \left\{ \sum_{\tau=1}^s A_{n\tau} e^{2\pi i(\mathbf{n} \cdot \mathbf{u}_{\tau})} \right\} e^{\frac{2\pi i}{a}(\mathbf{n} \cdot \mathbf{r})}, \quad (7)$$

where  $A_{n\tau}$  are the coefficients of the potential for a simple cubic lattice composed of atoms of a type denoted by  $\tau$ , and  $\mathbf{a} \cdot \mathbf{u}_{\tau}$  is the displacement of the atom  $\tau$  from the origin of the unit cell.

If  $\phi_{\tau}(\mathbf{r})$  be the potential due to a single atom of the type  $\tau$  we have by definition of the  $A_{n\tau}$

$$\sum_{\mathbf{m}} \phi_{\tau}(\mathbf{r} - \mathbf{m}\mathbf{a}) = \sum_{\mathbf{n}} A_{n\tau} e^{\frac{2\pi i}{a}(\mathbf{n} \cdot \mathbf{r})}$$

Hence

$$A_{n\tau} = \frac{1}{a^3} \int \phi_v(r) e^{-\frac{2\pi i}{a}(n \cdot r)} dv \quad (8)$$

This integral is just the well-known atomic scattering factor for electrons,<sup>†</sup> and if the charge distribution surrounding an atom be assumed spherically symmetrical, it may be put in the form

$$A_{n\tau} = \frac{e}{4\pi^2 n^2} (Z - f_n), \quad (9)$$

where  $f_n$  is the atomic scattering factor for X-rays taken at the value of its argument  $\frac{\sin \theta}{\lambda} = \frac{n}{2a}$  and  $Ze$  is the nuclear charge

If we write

$$\theta_\tau = 2\pi (n, u_\tau)$$

then, since the coefficients  $A_{n\tau}$  are real, we obtain by comparing (5) and (7)

$$|V_n|^2 = (\sum_\tau A_{n\tau} \cos \theta_\tau)^2 + (\sum_\tau A_{n\tau} \sin \theta_\tau)^2 \quad (10)$$

$\theta_\tau$  is just the phase difference between the origin of the unit cell, and the atom  $\tau$  when the crystal is reflecting X-rays from the planes  $(n_1, n_2, n_3)$  in the first order

Thus if the crystal structure and the atomic scattering factor are known from X-ray data it is always possible to obtain  $\Delta E_n$ , the discontinuity of the energy across any given plane in  $k$  space, to the accuracy given by the perturbation method. Although the absolute values of  $\Delta E_n$  obtained in this way may not be very accurate, one may expect to obtain satisfactory values of the relative energy gaps for the different planes

In the special case when the atoms forming the crystal have nearly the same atomic number, so that the atomic scattering factors may be taken to be equal, a simple approximate formula may be given for the energy gap across a given plane —

$$\Delta E_{n_1, n_2, n_3} = \frac{2e^2}{\pi^2 n^2 a} (Z - f_n) S_{n_1, n_2, n_3}, \quad (11)$$

where

$$S_{n_1, n_2, n_3} = \{(\sum_\tau \cos \theta_\tau)^2 + (\sum_\tau \sin \theta_\tau)^2\}^{\frac{1}{2}}$$

is the structure amplitude, which largely determines the relative intensities of the X-rays reflected by the crystal planes  $(n_1, n_2, n_3)$

<sup>†</sup> Mott and Massey, "Theory of Atomic Collisions," chap VII, Mott, 'Leipziger Vorträge,' p 63 (1930)

Even for those alloys for which there is a considerable difference between the atomic numbers of the constituent atoms, there will be a close similarity between the relative intensities of the X-rays reflected by the different sets of planes ( $n_1, n_2, n_3$ ) and the energy gaps across the corresponding planes in  $k$  space. The reason for this is that it is known that the X-ray reflection spectra of the three alloys Cu-Zn, Ag-Zn, Au-Zn are very similar, and that the relative intensities reflected from the different planes are principally determined by the values of  $(S_{n_1, n_2, n_3})^2$ . Hence, since for given  $n$  the difference between the values of  $(Z - f_n)$  for two different atoms is smaller than the difference between the values of  $f_n$ , it follows that the energy gaps will depend on  $S_{n_1, n_2, n_3}$  to an extent at least as great as the extent to which the X-ray reflections depend upon  $(S_{n_1, n_2, n_3})^2$ . Therefore, as the X-ray reflections for the three alloys are very similar, it follows that the energy gaps across the same planes in  $k$  space for the different alloys will also be approximately equal.

### 3 Alloys with the $\gamma$ -structure

Bradley and Thewlis† have made a detailed examination of the structure of the alloys Cu-Zn, Ag-Zn, and Au-Zn in the  $\gamma$ -phase, based upon the X-ray measurements made by Westgren and Phragmen‡. These measurements show that of the crystal planes with small indices only the set {411} and the set {330} give rise to strong X-ray reflections. We deduce, therefore, that the first planes in  $k$  space, outwards from the origin, across which the energy shows an appreciable discontinuity are those lying parallel to the sets {411} and {330}, and whose perpendicular distance from the origin is, in each case according to equation (4),  $3/2\sqrt{2}$ . A perspective drawing of the zone marked out in  $k$  space by these planes is given in fig. 2. The polyhedron has 36 faces and is very symmetrical. The ratio of its volume to that of the inscribed sphere is  $5\sqrt{2}/2\pi, \approx 1.12$ . The volume is equal to 45, and since there are  $2/3$  states per atom, the zone must contain 90 states per unit cell, and the inscribed sphere very nearly 80.

For the alloy Cu-Zn Westgren and Phragmen (*loc cit*) give for the value of the lattice constant  $a = 8.86 \text{ \AA}$ , therefore with  $n = \sqrt{18}$  one finds  $n/2a = 0.239 \cdot 10^8$ , the value of the atomic scattering factor  $f$  for Cu at  $(\sin \theta)/\lambda = 0.239 \cdot 10^8$  may be interpolated from the data given by James and Brindley§ and is found to be 19.9. Bradley and Thewlis (*loc cit*) give for the square

† 'Proc. Roy. Soc., A', vol. 112, p. 678 (1926).

‡ 'Phil. Mag.', vol. 50, p. 331 (1925).

§ 'Phil. Mag.', vol. 12, p. 81 (1931).

of the structure amplitude  $S^2_{411} = 31.7$  and  $S^2_{330} = 78.3$ . Thus using equation (11) we obtain the approximate values of the energy gaps across the planes (411) and (330) in  $k$  space

$$\Delta E_{411} = 0.93 \text{ electron volts}$$

$$\Delta E_{330} = 1.46 \text{ electron volts}$$

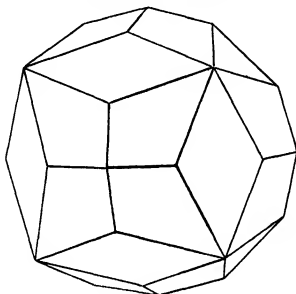


FIG. 2

These values are probably an over-estimate due to the approximations involved in the perturbation calculation which gives (6)

Within the  $\gamma$ -phase the formulae for the alloys Cu-Zn, Ag-Zn, Au-Zn are  $\text{Cu}_2\text{Zn}_8$ ,  $\text{Ag}_2\text{Zn}_8$ ,  $\text{Au}_2\text{Zn}_8$ , and there are 52 atoms per unit cell. If the number of loosely bound electrons contributed by each atom is given by the valency, i.e., Cu 1, Zn 2, Al 3, Sn 4, and so on, then throughout the whole set of alloys with the  $\gamma$ -structure there are as Hume-Rothery points out just 21 loosely bound electrons to every 13 atoms at a composition in the neighbourhood of the centre of the phase. Thus there are 84 electrons per unit cell, we have seen that the zone is nearly spherical, contains 90 possible states, and that the inscribed sphere contains 80, hence the surface of the Fermi distribution must lie at all points very close to the surface of discontinuity in the energy. This is the distinguishing feature of alloys with the  $\gamma$ -structure.

If the energy of the states associated with the points lying along a line in  $k$  space through the origin, and perpendicular to a plane of discontinuity, is

plotted against the distance from the origin, a curve of the form shown in fig 3 is obtained. The dotted line represents the energy of free electrons. Although from the present standpoint it is impossible to say anything about the actual value of the binding energy of the metal, it can be seen from fig 3 that the effects of the energy gaps, considered by themselves, will be to give an additional contribution to the negative energy, when the surface of the Fermi distribution lies just within or just touching the boundary of a zone in  $k$  space. Thus with the structure characteristic of the  $\gamma$ -phase we may expect the composition of the centre of the phase to correspond with somewhat less than 90 loosely

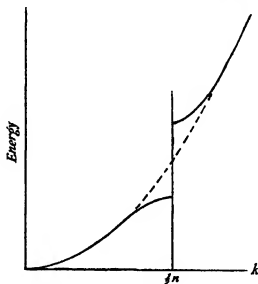


FIG 3

bound electrons per unit cell of 52 atoms. For, on account of the smallness of the energy gaps, 90 electrons per unit cell would cause an overlapping into the second zone, and the energy minimum must occur just before this happens. The Hume-Rothery rule gives 84 electrons.

#### 4 Diamagnetism

Let  $\kappa$  be the distance in  $k$  space measured normal to a plane across which the energy is discontinuous. For the discussion of the diamagnetism we require the value of  $\partial^2 E / \partial \kappa^2$ , see, for example, equation (12). It is easy to find the values of  $\partial^2 E / \partial \kappa^2$  at the origin, and by an elementary perturbation calculation, at either side of a plane of discontinuity. At the origin the value is simply

$\hbar^2/ma^2$ , which if the electrons were entirely free would be the value at all points in  $k$  space. The first approximation to the energies of states associated with points which lie close, on either side, to a plane of discontinuity,  $\pm e$ , for which  $k$  is nearly equal to  $\frac{1}{2}n$ , are given by the roots of the equation

$$\begin{vmatrix} \frac{\hbar^2}{2ma^2}|n-k|^2 + V_0 - E, & V_n \\ V_n^*, & \frac{\hbar^2}{2ma^2}|k|^2 + V_0 - E \end{vmatrix} = 0$$

Hence by use of (6) we obtain

$$\begin{aligned} \left(\frac{\partial^2 E}{\partial k^2}\right)_{k=n} &= \frac{\hbar^2}{ma^2} \left(1 - \frac{4E_{1n}^{(0)}}{\Delta E_n}\right) \\ \left(\frac{\partial^2 E}{\partial k^2}\right)_{k=0} &= \frac{\hbar^2}{ma^2} \left(1 + \frac{4E_{1n}^{(0)}}{\Delta E_n}\right), \end{aligned}$$

where  $E_{1n}^{(0)}$  is the energy of a free electron at the point  $\frac{1}{2}n$ , and  $\Delta E_n$  is the jump in the energy across the plane. Thus with the estimated values of  $\Delta E_n$  given for the alloy  $\text{Cu}_5\text{Zn}_3$  the magnitude of  $\partial^2 E/\partial k^2$  in the immediate neighbourhood of the planes {411} will be 37 times its value for free electrons, and in the neighbourhood of the planes {330} 24 times. This may easily be seen by reference to fig. 3.

Recently Pierls† has examined the diamagnetic susceptibilities of electrons in metals. He finds three terms which contribute to the susceptibility. For the most important of these terms he gives a formula which for metals at low temperatures may be written

$$\chi = -\frac{2m^2\mu^2}{3\hbar a} \iint \left(\frac{\partial E}{\partial n}\right)^{-1} \left\{ \frac{\partial^2 E}{\partial k_x^2} \frac{\partial^2 E}{\partial k_y^2} - \left(\frac{\partial^2 E}{\partial k_x \partial k_y}\right)^2 \right\} dS, \quad (12)$$

where the integration is over the surface of the Fermi distribution,  $m$  is the mass of the electron, and  $\mu$  its magnetic moment.  $\partial E/\partial n$  is the rate of change of  $E$  along the normal to the surface over which the integration is taken. This formula was obtained for a special case corresponding to the Bloch limiting case of tight binding. Since, however, the formula reduces for free electrons just to Landau's‡ formula it appears justifiable to use it in intermediate cases.

In general the magnitude of  $\frac{\partial^2 E}{\partial k_x^2} \frac{\partial^2 E}{\partial k_y^2}$  will be much greater than the magni-

† 'Z. Physik,' vol. 80, p. 763 (1933)

‡ 'Z. Physik,' vol. 64, p. 629 (1930)



tude of  $\left(\frac{\partial^2 E}{\partial k_x \partial k_y}\right)^2$  at all points on the surface over which the integration is taken. Hence, when the surface of the Fermi distribution lies close to a surface of energy discontinuity  $\chi$  may be expected to have a value many times greater than the value given by free electrons.

The experiments of Endo (*loc cit*) show that the alloys Cu-Zn and Cu-Sn have a remarkable maximum in the diamagnetic susceptibility for compositions which lie within the  $\gamma$ -phase. Fig. 4 is a reproduction of Endo's

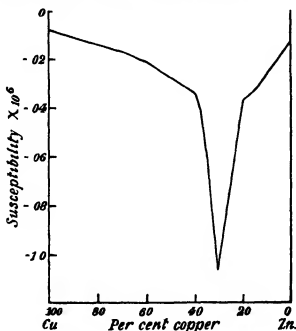


FIG. 4

curve showing the diamagnetic susceptibility as a function of the composition of the alloy Cu-Zn.

### Hall Effect

The following theoretical formula for the Hall coefficient has been obtained†

$$R = \frac{\alpha^2}{2e} \frac{\iint \left(\frac{\partial E}{\partial n}\right)^{-1} \left\{ \left(\frac{\partial E}{\partial k_x}\right)^2 \frac{\partial^2 E}{\partial k_y^2} - \frac{\partial E}{\partial k_x} \frac{\partial E}{\partial k_y} \frac{\partial^2 E}{\partial k_x \partial k_y} \right\} dS}{\left[ \iint \left(\frac{\partial E}{\partial n}\right)^{-1} \left(\frac{\partial E}{\partial k_x}\right)^2 dS \right]^2},$$

† Jones and Zener (in course of publication)

where in this case the integration is taken over that part of the surface of the Fermi distribution where it is not in contact with a surface of discontinuity

It can be seen that if the surface of the Fermi distribution lies in the main just within the boundary of the zone the Hall coefficient will be large and positive, i.e., of the same sign as for zinc, whilst if it lies just outside the boundary it will be large and negative

By slightly varying the composition of an alloy the number of loosely bound electrons may be varied slightly without changing the structure of the phase. For the  $\gamma$ -structures a small change in the number of loosely bound electrons would change the surface of the Fermi distribution from lying just within to just outside the boundary of the zone (cf fig 1). For these alloys therefore it may be expected that the Hall coefficient will change for small variations in the composition from large positive to large negative values as the composition varies in the direction of increasing number of loosely bound electrons. The experimental curves given by Evans and Richards (*loc cit*) do appear to behave in this way.

It is a pleasure to express my thanks to Dr C Zener and Professor N F Mott for discussions on various points.

### *Summary*

The zones of allowed energies for the loosely bound electrons in alloys with the  $\gamma$ -structure are examined. It is shown that the lowest group of energy levels is almost completely filled by the number of loosely bound electrons given by the Hume-Rothery rule, and a reason for the existence of this rule is discussed. In addition it is shown that large diamagnetic susceptibilities may be expected for alloys in this phase, and that the Hall coefficient should change from large positive to large negative values as the composition varies through the phase in the direction of increasing number of loosely bound electrons.

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*Some Experiments on the Production of Positive Electrons*

By J CHADWICK, F R S, P M S BLACKETT, F R S, and G P S OCCHIALINI

(Received February 10, 1934)

[PLATES 2-5]

§ 1 *Introduction*

This paper describes work carried out during the last year on the production of positive electrons, or positrons\* by various radiations. Although it is probable that the positrons found in conjunction with cosmic rays have their origin in the interactions of some penetrating radiation with matter, these experiments gave no direct information about their mode of origin. In order to throw light on this and to have a method of studying their properties more closely it was necessary to seek a means of producing positrons under easily controlled conditions.

Our first experiments directed to this end were those which showed that the radiation from beryllium bombarded by polonium  $\alpha$ -particles, a radiation consisting of  $\gamma$ -rays and neutrons, could eject positrons from lead. A preliminary notice of these results was given by us in a letter to 'Nature'†. Almost at the same time Meitner and Philipp‡ and Curie and Joliot§ reported the same effect.

We next proceeded to see whether positrons were produced by the high energy  $\gamma$ -rays emitted by thorium C'', and our first results were reported by Chadwick in the Bakerian lecture|| last year. By this time Curie and Joliot had shown that the positrons produced by the beryllium polonium radiation were mainly due to the  $\gamma$ -rays, rather than the neutrons, and first Anderson¶ then Curie and Joliot\*\* and Meitner and Philipp†† reported the production of positrons by the  $\gamma$ -rays of thorium C'''. It was pointed out by Blackett and

\* The name 'positron' for a particle of about electron mass and positive charge was suggested by Anderson, and it seems to be coming into general use. The negative electron will still be called, as usual, an electron.

† Chadwick, Blackett, and Occhialini, 'Nature,' vol 131, p 473 (1933).

‡ 'Naturwiss,' vol 21, p 286 (1933).

§ 'C R Acad Sci Paris,' vol 196, p 1105 (1933).

|| 'Proc Roy Soc,' A, vol 142, p 1 (1933).

¶ 'Science,' vol 77, p 432 (1933).

\*\* 'C R Acad Sci Paris,' vol 196, p 1581 (1933).

†† 'Naturwiss,' vol 24, p 468 (1933).

Occhialini\* that the Dirac theory of the electron led to the view that a pair of electrons of equal mass and opposite sign may be produced when  $\gamma$ -rays of high energy are absorbed in matter. If  $m$  is the mass of either electron then the combined kinetic energies of the particles will be

$$E_2 = h\nu - 2mc^2 \quad (1)$$

and this will also be the maximum kinetic energy which the positron can acquire †. Since the mass of the electron is equivalent to an energy of 511,000 electron volts the total kinetic energy of the pair (or the maximum energy of the positron) produced in this way by the thorium C''  $\gamma$ -ray of  $h\nu = 2.62 \times 10^6$  e volts will be  $1.60 \times 10^6$  e volts. Anderson and Neddermeyer‡ have, in fact, measured the energies of the two particles for several pairs and they have found that their total kinetic energy does not appreciably exceed this value, and Curie and Johot and Meitner and Philipp have measured single positrons with energies up to about this value. The preliminary results of our own measurements, quoted in the Bakerian lecture, also led to the same conclusion. A survey of these results has recently been given by Blackett §.

It is clearly of the greatest importance to establish the above relation thoroughly, for upon it largely depends the experimental evidence that the positron is created, simultaneously with an electron, by some interaction of a  $\gamma$ -ray with matter, and, moreover a value for the mass of the positron can be deduced from it. The experiments were therefore continued until about 4000 tracks of electrons and about 400 tracks of positrons had been obtained, giving a body of evidence sufficient to justify quantitative conclusions.

## § 2 *Experiments with the $\gamma$ -rays of Thorium Active Deposit*

The expansion chamber used in these experiments was the one made by Blackett and Occhialini for the study of cosmic rays. The chamber had an internal diameter of 1.3 cm. and the piston travelled horizontally. The chamber was filled with oxygen, to a pressure of about 1.7 atmospheres. The chamber was placed between two coils which served to establish a magnetic field with its lines parallel to the axis of the piston. The strength of the field was in most experiments about 800 gauss, in others, 400 and 600 gauss. The tracks

\* 'Proc Roy Soc,' A, vol 139, p 699 (1933)

† It will acquire the maximum energy when the electron, at the end of the process, is free and at rest.

‡ 'Phys. Rev.', vol 43, p 1034 (1933)

§ 'Nature,' vol 132, p 917 (1933)

were photographed by means of two cameras, one with its axis along the axis of the piston, the other at an angle of  $20^\circ$  to this

In the first series of experiments, the source of thorium active deposit, of about 1/50 mg  $\gamma$ -ray activity, was placed outside the chamber in a lead cylinder. The  $\gamma$ -radiations, filtered through 1 cm of lead, passed through the glass walls of the chamber and fell on a target of lead, 2 cm square and 2 mm thick, attached to the inside wall.

In order to avoid any possibility that a negative electron moving towards the lead target should be counted as a positron, a thin metal plate, of aluminium or of copper, was placed across the middle of the chamber so as to intercept the path of the particles, following the arrangement used originally by Anderson. The particle must lose energy in passing through the plate and its track must therefore show a greater curvature on the emergent side than on the incident side. The direction of motion, and therefore the sign of the charge of the particle, can then be definitely known. In this series some 1200 tracks were obtained about 50 of which could with certainty be ascribed to particles carrying a positive charge. The ionizing power of these particles and the loss of energy in passing through the plate were both consistent with the view that the particles had a mass about the same as that of the negative electron. Occasionally positive electrons were observed from the glass walls of the chamber, and also from the plate of copper across the chamber.

In a second series of experiments the source of thorium active deposit was placed inside the chamber. The deposit was obtained on a wire about 5 mm long and this was surrounded by a closed cylinder of lead of 3 mm wall. The lead cylinder was fixed to the top plate of the expansion chamber. With this arrangement, the source occupied a very small volume and the chance that a negative electron produced in a remote part of the chamber should be bent so as to hit the lead cylinder was very small. It seemed no longer necessary to place a metal plate across the chamber in order to identify the sign of the charge in the particle, and the plate was therefore omitted, as its presence interfered with the accurate measurement of the curvature of the tracks.

About 700 pairs of photographs were taken in this series of experiments giving more than 4000 tracks of negative and positive electrons.

The curvature of the tracks was measured by superimposing over the image of a track a transparent scale with circles of different radii marked on it. This method is not a very accurate one, but it does serve to avoid errors due to any departure of the track from a true circle such as might be caused by small nuclear deflections. Experience has shown that such deflections are difficult

to recognize and may cause serious errors. This method enabled the energy of a particle to be determined with a probable error of about 4%. A further source of error was present because no attempt was made to correct for the actual position of the track in the chamber. On this account the true magnification of the different tracks varied as much as 3% from the mean. Nor, when calculating the energy of the particle from the track curvature, was any correction made for the fact that a track might not lie exactly in the plane normal to the magnetic field. This error was, however, small since only those tracks were selected for measurement the planes of which, as revealed by comparison of the two stereoscopic photographs, were within a small angle to the plane normal to the field. Clearly it would have been desirable to improve the accuracy of the measurements by taking all these sources of error into account, but we considered that the great labour was not justified for a preliminary survey of the phenomena, and we preferred to select from the photographs those tracks which were most suitable for measurement.

The results of the measurements of the curvatures of the tracks are shown in fig. 1. Curve 1 gives the data for all the positrons with energies greater than  $0.5 \times 10^6$  volts which were suitable for accurate measurement, and curve 2 for the electrons with energies above  $1 \times 10^6$  volts, selected from about one-tenth of all the photographs. In each curve the number of tracks within an energy range of 200,000 volts is plotted against the mean of the limit of the energy range. It must be pointed out that these curves do not represent the energy distributions of the electrons as they are ejected by the  $\gamma$ -radiation, for the thickness of the lead absorber in which the electrons arise is greater than the maximum range of the fastest particles. Thus electrons of all energies from zero to the maximum must be present, whatever the original distribution may be. No detailed interpretation of the curves of energy distribution is possible owing to the complexity of the experimental conditions.

The  $\gamma$ -radiation emitted by thorium active deposit consists of (1) the soft radiation of thorium B, with energies up to 300,000 volts, (2) the radiation from thorium (C + C'), mostly of energies below about 700,000 volts but with two weak lines of 1.6 and 1.8 million volts, (3) those of thorium C'', some of low energy up to 600,000 volts and the strong line of 2.62 million volts. If we restrict our examination of the electrons to those of high energy then only the  $\gamma$ -rays of 1.6 and 1.8 million volts, associated with the thorium C - C' transformation, and that of 2.62 million volts, arising from the Th C'' - Pb transformation, are concerned. The first two rays are of weak intensity and, although twice as many atoms transform from Th C to C' as from Th C'' to Pb,

about 70% of the quanta of high energy must be contributed by the line of 2.62 million volts. The end portions of the curves of fig. 1 must then be due mainly to the action of this  $\gamma$ -ray. It seems reasonable for a first approximation to assume that the maximum energy of the particles is given by the intercept on the energy axis of the steep portion of the curve, as drawn in fig. 1.

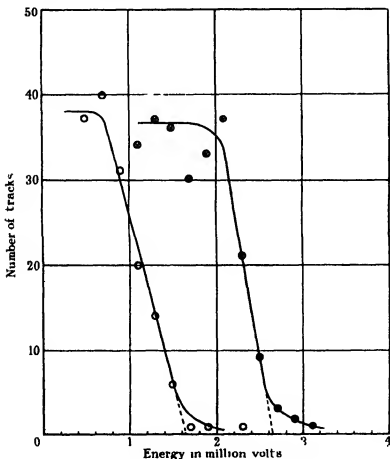


FIG. 1 — O Positrons, ⊗ electrons

The end point as thus defined will be too high by a quantity somewhat greater than the mean error of measurement \*

\* The effect of the errors of measurement in the track curvatures is to increase the slope of the end part of the curves of fig. 1. Suppose, for example, that the energy distribution is uniform up to a maximum energy  $E_0$  (see fig. 2). If each energy determination is subject

The measured end points for the positron and for the electron curves are  $1.63 \times 10^6$  volts and  $2.62 \times 10^6$  volts respectively, with estimated probable errors of about  $1\frac{1}{2}\%$ . The calculation given in the note shows that these values should be about 5% high. We thus get for the corrected values of the maximum energies of the positron and electron the values

$$E_2 = (1.55 \pm 0.03) \times 10^6 \text{ volts}$$

$$E_1 = (2.49 \pm 0.04) \times 10^6 \text{ volts}$$

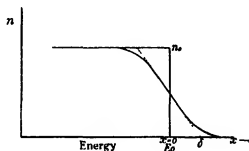


FIG. 2

The value found for  $E_2$  is sufficiently close to the value of  $1.60 \times 10^6$  volts given by equation (1) above to provide very strong support to the hypothesis that a  $\gamma$ -ray produces a negative electron at the same time as a positive electron and that the two have equal masses.

It is conceivable, however, that in contradiction to Dirac's theory the mass  $m_2$  of the positron is not exactly equal to the mass of the electron. If we rewrite (1) in the form

$$m_2 c^2 = h\nu - E_2 - m_1 c^2, \quad (2)$$

to an error, the probability of which is given by the Gaussian error law, then the true distribution curve ( $n = n_0$  for  $x < 0$  and  $n = 0$  for  $x > 0$ ) is transformed into the curve

$$n(x) = \frac{n_0 h}{\sqrt{\pi}} \int_{y=-\infty}^{y=0} e^{-h^2(y-x)^2} dy$$

Since  $n(x) = \frac{1}{2}n_0$  when  $x = 0$ , the value of the energy at which the number of particles passes through half value gives the true end point  $E_0$ . The tangent to the distribution curve at this midpoint cuts the  $x$  axis at a value  $\delta$  given by  $\delta = \sqrt{\frac{\pi}{2}} \mu = 1.25 \mu$ , where  $\mu$  is the R.M.S. error of the observations. For this special case then, the quantity  $\delta$  must be subtracted from the observed end point to give the true value of  $E_0$ . Since the R.M.S. error of the energy determination was estimated at about 4%, the end point should be about 5% too high.





FIG. 3



FIG. 4



FIG. 5



FIG. 6



FIG. 7



FIG. 8



FIG. 9



FIG 10



FIG 11

and introduce the observed value of  $E_2$  and the known value of  $h\nu$  we obtain

$$m_2 = (1.10 \pm 0.06) m_1$$

A preferable procedure, which reduces the error introduced by any small constant error in the absolute values of the measured energies, is to use the observed value of  $E_1$  rather than the value of  $h\nu$ . The majority of the observed electrons are recoil electrons, which have a maximum energy of  $2.39 \times 10^6$  volts. The photo-electrons are probably about 10% of the recoil electrons and mostly come from the K shell, the contributions of the L and other shells being quite small. It is very probable therefore that the end point of the electron curve as we have defined it will correspond to the maximum energy of the photo electrons from the K shell, that is,

$$E_1 = h\nu - \omega_K, \quad (3)$$

where  $\omega_K$  is the binding energy of the K shell.

From (2) and (3) we get

$$m_2 c^2 = E_1 - E_2 + \omega_K = m_1 c^2$$

Since  $\omega_K = 0.09 \times 10^6$  volts, we obtain

$$m_2 = (1.02 \pm 0.10) m_1$$

This calculation gives probably the most accurate determination of the mass of the positron which is yet available.

It will be noticed that both curves of fig. 1 show a tail, particularly marked for the electron curve. Such a tail is due in part to errors of measurement, possibly caused by unrecognized deflections in the tracks, but it may also be partly due to the presence of a small amount of  $\gamma$ -radiation of energy greater than  $2.62 \times 10^6$  e. volts.\* The presence of a few cosmic ray tracks passing through the source may perhaps contribute to these tails, the vertical type of cloud chamber, such as we used, is sensitive to cosmic radiation.

### § 3 Yield of Positive Electrons

In the first series of experiments the number of positive electrons obtained in 360 pairs of photographs was 50, compared with 1200 negative electrons. In the counting of the positive electrons an over strict criterion was adopted

\* Dr. Ellis informs us that the nuclear level system proposed by Mott and himself for the thorium C'  $\gamma$ -rays suggests a radiation of  $h\nu = 3.2 \times 10^6$  e. volts.

The yield of positive electrons, about 4% of the negatives, obtained from this series is certainly much too low

In the second series, in which the source of  $\gamma$ -rays was inside the expansion chamber, all tracks which started from the lead cylinder and which showed a positive curvature were counted as positrons. A corresponding criterion was taken for the negative electrons. Two counts were made, in the first all particles of energy greater than about 200,000 volts were counted, and in the second only those of energy greater than 400,000 volts. The first gave 323 tracks of positrons and 3563 tracks of negative electrons. The ratio of positive to negative electrons is thus about 9% and may be slightly greater, for some positive tracks which could not be traced to their origin in both photographs of the stereoscopic pair were rejected as doubtful. The second count gave a similar proportion.

The actual production of positrons in the lead absorber must be greater than this observed ratio, owing to the fact that the energy of the positives is less than that of the negatives and a greater fraction will be unable to escape from the absorber. A rough estimate of the correction to be applied for this absorption can be made in the following way.

Consider the production of secondary electrons in a thick layer of matter by a homogeneous  $\gamma$ -ray, and assume that all the secondary electrons are emitted directly forwards with the same energy  $E_0$ . Then the number which emerge from the absorber with energies greater than  $E_1$  will be proportional to  $\alpha(R_0 - R_1)$ , where  $\alpha dx$  is the chance of production of a secondary electron in a layer of thickness  $dx$  and  $R_0$  and  $R_1$  are the ranges in the absorbing material of electrons of energy  $E_0$  and  $E_1$ . Thus in this case the ratio of the observed number  $N'$  of positrons to the number  $N$  of electrons both with energies greater than  $E_1$ , will be

$$\frac{N'}{N} = \frac{\alpha'}{\alpha} \frac{R'_0 - R'_1}{R_0 - R_1},$$

where  $\alpha'$ ,  $R'_0$ ,  $R'_1$  refer to the positrons. It is here assumed that positive electrons have the same range as negative electrons of the same energy. From the known data on the ranges of  $\beta$ -particles we find the following data -

$$\begin{array}{ll} E_0 = 2.39 \times 10^6 \text{ volts}^* & R_0 = 1.12 \text{ gm./sq. cm.} \\ E'_0 = 1.60 \times 10^6 \text{ volts} & R'_0 = 0.67 \text{ gm./sq. cm.} \\ E_1 = 0.2 \times 10^6 \text{ volts} & R_1 = 0.04 \text{ gm./sq. cm.} \end{array}$$

\* One must take here the values for the recoil electrons since these are in great majority

Taking the observations in which all particles with energies greater than 200,000 volts were counted,  $N'/N$  was 9%, whence  $\alpha'/\alpha = 15\%$ . This calculation is very rough for several factors have been neglected. For example, no account has been taken of the angular distribution of the particles. It is possible that the positrons are emitted mainly within small angles of the direction of the  $\gamma$ -ray, and the above correction for escape will then be over-estimated. It seems likely that the probability of production of a positron in the lead absorber is from 12 to 15% of the probability of the production of a recoil electron.

Now the  $\gamma$ -radiation passing through the lead consists, as we have already described, mostly of  $\gamma$ -rays of low energy with three  $\gamma$ -rays of energy 1.6, 1.8, and 2.62 million volts. If the Dirac theory of the production of positive electrons is correct then the positive electrons can be due only to  $\gamma$ -rays of energy greater than  $1.02 \times 10^6$  e volts. Further, it seems probable, both from theory and from such experimental information as is available, that the probability of production increases rapidly with the energy of the  $\gamma$ -ray. Thus the positrons observed in our experiments must be due almost entirely to the  $\gamma$ -ray of 2.62 million volts, while the other radiations present take part in the production of the recoil electrons. It is only possible to make a very rough estimate of the proportion of the negative electrons which are produced by the  $\gamma$ -ray of 2.62 million volts, it seems likely that this fraction is between 0.5 and 0.7, under the conditions of our experiments. We thus arrive at the conclusion that the probability of the production of a positron by the  $\gamma$ -ray of 2.62 million volts in lead may be as high as 20 to 30% of the probability of the production of an electron by the normal processes of scattering and photoelectric absorption.

#### § 4 Experiments with a $\beta$ -ray Source

In some experiments a thin aluminium wire coated with thorium active deposit was enclosed in a glass tube, about 0.1 mm thick, and placed in the centre of the expansion chamber. With this arrangement the majority of the  $\beta$ -rays emitted by the active deposit could escape through the glass tube. The source was of much weaker activity than in the previous experiments, about 15 electron tracks being observed on the average in each expansion. A few tracks of positrons were also observed, in all 24 tracks among 2000 tracks due to electrons. The energy of the positrons was never greater than  $1.4 \times 10^6$  volts. A rough calculation based on the results of the previous sections shows that it is unlikely that these positrons could be due to the action of

$\gamma$ -rays in the metal of the wire or in the glass walls of the tube. Moreover, in some experiments the active deposit was obtained on a platinum wire instead of aluminium. If the positrons were produced in the wire, a much greater proportion should have been found in this case, but the proportion remained sensibly the same. Some photographs were taken in which a thin platinum cylinder was placed round the source tube to test whether the  $\beta$ -particles could produce positrons in passing through the platinum. No increase in the proportion of positives could be noticed. It seems probable therefore that these positive electrons have their origin in the radioactive atoms themselves, possibly by an internal conversion of a  $\gamma$ -ray or perhaps by an interaction of a  $\beta$ -particle in its escape through the nuclear field.

It may be pointed out that these results indicate that a bare source of thorium active deposit provides the most powerful source of positive electrons which is yet known. This is indeed evident from the experiments of Thibaud\* who has succeeded in obtaining from a radiothorium source a pencil of positive electrons sufficiently intense to give a good trace on a photographic plate, and to offer the possibility of measuring the ratio  $e/m$  of the charge to the mass.

### § 5 *Experiments with the Radiations Excited in Beryllium, Boron, and Fluorine*

The arrangement of the source and target in these experiments was similar to that in the first series with the thorium  $\gamma$  rays. A capsule containing a polonium source and a piece of beryllium was placed outside the expansion chamber close to the wall, and on the inside wall was fixed a target of lead, 2.5 cm square and 2 mm thick. The lead target was thus exposed to the  $\gamma$ -rays and neutrons liberated from the beryllium under the bombardment of the  $\alpha$ -particles from the polonium, and also to the  $\gamma$ -rays emitted by the polonium itself. Since the  $\gamma$  rays of polonium have a maximum energy of only 1 million volts, no confusion can arise if observations are restricted to electrons of energy greater than this amount. This fact was confirmed in control experiments in which the polonium alone was used as a source, and in these experiments no positive electrons were observed †. In most photographs a metal plate was placed across the middle of the expansion to intercept the paths of the positive electrons and to show unambiguously their direction of motion. In all the photographs with a lead target 203 negative electrons were

\* 'Nature,' vol. 132, p. 490 (1933).

† See also Meitner and Philipp, *loc. cit.*



obtained and 66 positive electrons, a proportion of the latter of about 32%. In some experiments the lead target was removed so that the electrons observed came from the glass wall of the chamber. In this case the proportion of positrons was much less, being only of the order of 4%.

The measurements of the curvature of the tracks showed that the end point of the positron curve was at about  $4 \times 10^6$  volts, and the end point of the electron curve at about  $5 \times 10^6$  volts. These results are similar to those already published by Curie and Johot\*. They are consistent with the view that the positrons are produced by a  $\gamma$ -ray of about  $5 \times 10^6$  e volts†. Some electron tracks, 10 in all were observed which had energies greater than  $6 \times 10^6$  volts but their curvatures were too small for accurate measurement. The evidence is too meagre to decide whether these tracks are due to a  $\gamma$ -radiation of high energy emitted from the beryllium or whether they are cosmic rays passing accidentally through the lead target.

Two sets of observations were made in which the beryllium in the source capsule was replaced first by boron then by calcium fluoride. With boron 11 positrons and 27 electrons were observed in 122 photographs a proportion of about 40%. The radiation from boron bombarded by polonium  $\alpha$ -particles consists partly of neutrons and partly of  $\gamma$ -radiation, the maximum energy of which appears to be about  $3 \times 10^6$  volts‡. The energy of the radiation is thus not very different from that of the strong line of the thorium radiation, and we should expect the efficiency of production of positrons in the two cases to be about the same. It seems very probable, therefore, that some of the positive electrons observed from boron must be attributed to the action of neutrons. This conclusion is supported by the observations made with fluorine, in which 12 positrons and 31 electrons were found in 139 photographs. As far as is known no  $\gamma$ -radiation is emitted from fluorine bombarded by polonium  $\alpha$ -particles with energy greater than about 1 million electron volts. It appears then that the positrons observed in this case must also be attributed to the action of neutrons. It is probable that the production of positrons by neutrons is an indirect effect. It is known that the passage of neutrons through lead is accompanied by the emission of  $\gamma$ -radiation, due perhaps to an excitation or a disintegration of the lead nucleus in a neutron collision. This  $\gamma$ -radiation, if of sufficient energy, will in its absorption in the lead produce positive electrons, possibly also by a type of internal conversion in the escape from the excited nucleus.

\* 'J Phys. Rad.', vol. 8, p. 494 (1933).

† Cf. Becker and Bothe, 'Z. Physik' vol. 70, p. 421 (1932).

‡ Cf. Becker and Bothe, *loc. cit.*

### § 6 Conclusions

It has been shown that when  $\gamma$ -rays of high energy are absorbed in matter positive electrons as well as negative electrons are ejected. The observations of the energies of the positrons ejected by the  $\gamma$ -rays of thorium C'' are consistent with the view that an electron is always produced at the same time as a positron. This is confirmed to some degree by the observation that occasionally pairs of tracks occur, one positively curved in the magnetic field, the other negatively, which are almost certainly due to the simultaneous production of a pair of electrons of opposite sign (see figs 5, 6, Plate 3). This event was, however, seldom observed. Presumably, under the conditions of our experiments in which the particles originated in a thick lead absorber, one or other of a pair was generally absorbed or scattered before emerging from the target. On the hypothesis that an electron is always produced at the same time as a positron the measurements lead to the result that the mass of the positron is the same as that of the electron.

It has been tacitly assumed that the pair of particles is produced in the atom rather than inside the nucleus. This view is supported by a consideration of the frequency of production of the positrons. In the case of beryllium radiation passing through lead, one pair is produced for rather less than two electrons produced by the normal scattering process. Assuming for this radiation the value of the scattering coefficient given by the Klein-Nishina formula for  $h\nu = 5 \times 10^6$  e. volts, we find that the effective cross section for the production of a positron is rather more than  $3 \times 10^{-24}$  cm<sup>2</sup>. This is distinctly larger than the area of cross-section of the lead nucleus.

Again, the results of § 3 gave a ratio of about 25% for the absorption of the thorium C''  $\gamma$ -rays by the production of positrons to the absorption by normal processes. From the total absorption coefficient of this radiation in lead,  $0.46$  cm<sup>-1</sup>, we obtain for the area of cross-section of the lead atom for the production of a positron the value  $2.8 \times 10^{-24}$  cm<sup>2</sup>. Recently Heitler and Sauter\* have calculated the cross section for the production of positrons from Dirac's theory. They find a value of  $2.6 \times 10^{-24}$  cm<sup>2</sup> for the case considered here. The good agreement, though to some extent fortuitous on account of the corrections applied to the experimental results, gives strong support to the validity of the theoretical calculations, particularly to the assumption that the positrons produced by the interaction of  $\gamma$ -rays with matter have their origin (mainly at least) in the field around the nucleus rather than inside it.

\* 'Natura', vol 132, p 892 (1933).

It may be that positrons produced by other agencies may arise from the nucleus itself, Curie and Joliot have found that positrons are produced when aluminium and boron are bombarded by  $\alpha$ -particles, and they suggest that these are ejected from the disintegrating nucleus

A necessary consequence of the production of the positron is its subsequent disappearance, for general experience shows that the positron has no permanent existence. If a quantum can interact with an atom to produce a pair of electrons of opposite sign, then the reverse process, in which a positron and an electron combine under the influence of an atom to produce radiation must also be possible. The positron cannot exist for more than a very short time in matter of ordinary density. We might then expect to observe the annihilation of a positron by the sudden disappearance of its track in the cloud chamber. This event has unfortunately not yet been observed with certainty. A few photographs have been obtained in which it appears that the positron has disappeared when still possessing a large amount of energy but it is not possible to be quite certain that the track has not merely passed out of the field of view or out of the illumination. Examples of this disappearance are shown in fig. 6, Plate 3, and fig. 10, Plate 5.

It was suggested by Blackett and Occhialini that the creation of positrons might account for the 'anomalous' absorption of high energy  $\gamma$ -rays, and that the radiations arising from the annihilation of the positrons might correspond to the secondary radiations which accompany the anomalous absorption. This question has been discussed at some length in a recent paper by Gray and Tarrant\*. They come to the conclusion that the creation and disappearance of positrons cannot account for all their experimental results, more particularly for those on the secondary radiations. Here we wish only to point out that the results of § 3 show that the extra absorption due to the production of positrons by the  $\gamma$ -ray of thorium C'' is just sufficient to account for the anomalous absorption. It may be that we have over-estimated the corrections to be applied to the observations and that the creation of positrons is not so frequent as we have suggested, but at least a large part of the anomalous absorption, if not all, must be attributed to this process.

We take this opportunity of thanking Mr. Ouellet for his assistance on many occasions during the course of the work. One of us (G. P. S. O.) is indebted to the Italian Council of National Research for a grant.

\* 'Proc. Roy. Soc.,' A, vol. 143, p. 706 (1934).

*Summary*

The emission of positive electrons has been observed under different experimental conditions (1) from a lead target exposed to the  $\gamma$ -rays of thorium active deposit, (2) directly from a source of thorium active deposit, and (3) from a lead target exposed to the radiations ( $\gamma$ -rays and neutrons) emitted by beryllium, boron, and fluorine when bombarded by polonium  $\alpha$ -particles. The measurements of the energies of the positrons ejected from lead by the thorium  $\gamma$ -rays support the view that a positron and an electron are produced simultaneously by the interaction of a  $\gamma$ -ray and an atom, and that the mass of the positron is the same as that of the electron. The positron and electron are probably created in the electric field outside, rather than inside, the nucleus. The observations show that when  $\gamma$ -rays of high frequency pass through lead an appreciable fraction (about one-fifth for a  $\gamma$ -ray of  $h\nu = 2.6 \times 10^6$  volts) of the energy absorbed is used in this process of creating a positron and an electron.

## DESCRIPTION OF PLATES.

The magnification is approximately the same in all photographs, and about 0.8

## PLATE 2

FIGS. 3 and 4.—The source was thorium active deposit inside a lead cylinder of 3 mm. wall. Magnetic field 740 gauss. Negative particles travelling to the right are bent downwards. On both photographs tracks of positrons, ejected from the lead by  $\gamma$  rays, can be seen. The  $\alpha$  track in fig. 4 is due to contamination.

## PLATE 3

FIG. 5.—Source of radiation, calcium fluoride bombarded by polonium  $\alpha$  particles, outside the chamber. The photograph shows a pair of tracks from the lead target, the upper track is due to an electron, the lower to a positron. Both particles have energies of about half a million volts.

FIG. 6.—Source of radiation, beryllium + polonium. The positron of this pair has an energy of  $2.8 \times 10^6$  volts. The track does not pass through the aluminium plate (0.90 mm. thick). The positron may have been annihilated while still possessing great energy, or it may merely have been deflected in such a way that it did not emerge from the plate.

FIG. 7.—Source was thorium active deposit in a thin glass tube, so that the  $\beta$ -rays could emerge. Two sharp  $\beta$ -ray tracks are seen, and one positron track travelling downwards.

## PLATE 4

In all four photographs, figs. 8–11, the source of radiation was beryllium + polonium.

FIG. 8.—An electron of energy  $3.9 \times 10^6$  volts passes through the aluminium plate, 0.33 mm. thick.

FIG 9 —A positron passes through an aluminium plate, 0.33 mm. thick, its energy on the upper side is  $1.02 \times 10^6$  volts, on the lower side  $0.7 \times 10^6$  volts

PLATE 5

FIG 10 —A positron track ends suddenly in the gas. The positron has probably been destroyed, but it is possible that it has merely passed out of the illumination

FIG 11 —A positron passes through a copper plate, 0.25 mm. thick, its energy on the upper side is  $3.2 \times 10^6$  volts, on the lower side  $2.6 \times 10^6$  volts



*The Influence of Pressure upon the Flame Spectra of Hydrogen and Carbonic Oxide*

By WILLIAM A BONE, D Sc, F R S and F G LAMONT, M Sc

(Received November 15, 1933)

[PLATES 6-10]

Some time ago, as a new development in high-pressure technique, apparatus was installed in our laboratories for the study of continuous flames at high pressures, and recently it has been adapted for spectrographic work. The present paper deals with observations on the influence of increasing pressure up to 100 atmospheres on the flame-spectra of hydrogen and carbonic oxide, and various mixtures thereof, burning in oxygen.

*Experimental*

*Apparatus* - Experience with the apparatus employed in previous experiments on the formation of nitric oxide in continuous high-pressure flames of carbonic oxide in oxygen-nitrogen atmospheres\* suggested certain improvements which led to the new design shown in elevation and section in figs 1 and 2. It consists of a central combustion chamber  $4\frac{1}{2}$  inches internal diameter and 1 inch deep enclosed by a massive ring AA, 12 inches diameter to which are bolted two end plates B and C  $3\frac{1}{2}$  inches thick. The ring carries the burner, the two inlet connections D and E and two quartz window plugs F and G. Each of the cover plates also carries a quartz window plug, H and K, and an outlet connection through which the products of combustion may be led away. By employing this method of construction it was possible to reduce the distance between the burner and the quartz windows H and K to less than  $\frac{1}{8}$  inch with a corresponding reduction in the absorption effects of the compressed gases in the chamber.

The cover plates are secured by six bolts of "Vibrac" steel and are designed to withstand a working stress of 10 tons each, which corresponds, after allowing for that due to tightening, to an internal pressure in the chamber of 300 atmospheres. The joints are made with Klungerite washers. The body, cover plates, and plugs were constructed of stainless steel (E S C Immaculate 3) to resist the corrosive effects of the hot gases from the flame.

\* Newitt and Lamont, 'Proc Roy Soc,' A, vol 139, p 83 (1933)

The system of valves, gauges, and connections supplying the compressed gases from the respective storage cylinders (100 to 120 atmospheres) to the combustion chamber was similar to those previously employed,\* and included two bronze tubes in series, the one packed with activated charcoal for the removal of any traces of iron carbonyl from the compressed carbonic oxide, and the other with purified phosphoric anhydride for drying purposes. Also, a large capacity "trap" was inserted in the outlet pipe of the combustion

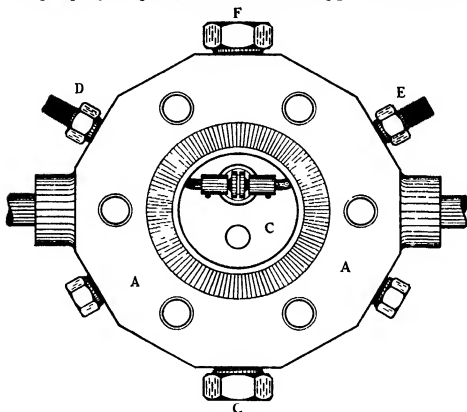


Fig 1

chamber to prevent the release valve becoming blocked by condensed water in experiments with hydrogen flames

The burner was on the "opposed-jet" principle employed in previous work. In this case, however, in lieu of the slotted tube surrounding the jets, each of them was fitted with a silica washer, 1 inch diameter and  $\frac{1}{8}$  inch thick, by means of which the flame could be maintained more accurately in the form of

\* *Loc cit.*, p. 86.

a vertical disc mid-way between the jets without touching either of them. And by carefully focussing a slightly enlarged image of the flame on the slot of the Hilger E<sub>2</sub> quartz-spectrograph employed it was possible entirely to prevent any continuous "black-body" radiation from the hot silica discs reaching

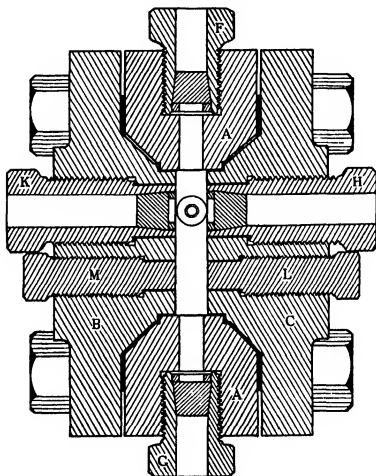


FIG 2

the plate on which the flame spectra were photographed, the images of the discs falling quite clear of the actual slit

*Procedure*—The experimental procedure consisted in maintaining steady flames of carbonic oxide, hydrogen, or known mixtures thereof, in an atmosphere of oxygen at each of the selected pressures (1, 10, 30, and 100 atmospheres), and after focussing them on the slit of the spectrograph in exposing its photographic plate for a period such that approximately equal amounts of gas were



burnt during each exposure. The exposure diminished from  $2\frac{1}{2}$  hours for flames at atmospheric pressure down to 2 minutes only for those at 100 atmospheres, the flames being maintained, as nearly as could be, of constant size.

Oxygen was employed as the supporter of combustion so as to eliminate any effects of oxides of nitrogen, it was always maintained in excess, namely in approximately equal volumetric ratio to the gas burnt. The combustible consisted first of pure dry carbonic oxide,\* and thence through a graded series of CO- $H_2$  mixtures, namely, 93CO/7 $H_2$ , 78CO/22 $H_2$ , 50CO/50 $H_2$ , 22CO/78 $H_2$ , and 7CO/93 $H_2$  finally to pure hydrogen itself, because it was desired to study not merely the influence of pressure upon the flame spectra of hydrogen and carbonic oxide, but also, at each pressure, the effects of successive  $H_2$ -additions to the CO-flame, and thus extending Weston's work to higher pressures †

*Weston's Observations*—It may be recalled how Weston's spectrogram of an undried CO-flame maintained at atmospheric pressure had shown, besides two groups of "steam" ( $H_2O$ , OH) bands, a banded radiation extending from 5000 Å in the visible region to 2200 Å in the ultra-violet, on which a continuous spectrum was superposed. Fowler and Gaydon‡ have recently concluded (in general agreement with and amplification of what was said in Weston's paper) that the banded radiation referred to is due to the direct combination of excited neutral CO molecules with neutral  $O_2$  molecules without dissociation into atoms of carbon and oxygen or molecules of carbon. The superimposed continuous spectrum which Weston showed is also associated with the direct oxidation of carbonic oxide in the flame, is due to the emission of unquantized radiation during the combustion.

Weston also found that as the carbonic oxide in the burning gas was progressively replaced by hydrogen, both the banded and the continuous parts of the burning CO-spectrum rapidly faded away, until with an equimolecular CO- $H_2$  mixture they had nearly all disappeared leaving only the "steam lines" visible in the spectrum. What were then termed "steam lines" are now known to be OH bands, and in what follows will be so described.

*Repetition of Weston's Experiments*—As it was thought advisable, for comparative purposes, to include in the present series a repetition of Weston's observations with flames at atmospheric pressure, especially as we contemplated burning much larger quantities of gas, and were using a spectrograph of greater analysing power, than he did

\* Dried by insertion of a bronze tube (10 inches long) filled with purified phosphoric anhydride.

† 'Proc Roy Soc,' A, vol 109, pp. 523-524 (1925)

‡ 'Proc Roy Soc,' A, vol 142, p 389 (1933)

Accordingly in fig. 3, Plate 6, are reproduced our spectrograms for CO-H<sub>2</sub> flames in oxygen at atmospheric pressure. The CO flame spectrogram shows a continuous radiation extending to *circa* 2400 Å, and superposed upon (i) the banded CO-O<sub>2</sub> radiation previously observed by Weston, and (ii) the HO-band at *circa* 3064 Å, the intensity of which is relatively so weak that only the lines near the head are visible. On progressively adding hydrogen to the CO-flame, the continuous radiation rapidly diminished and HO bands became more pronounced until with a pure H<sub>2</sub>-flame four of them (with heads at 2608, 2811, 3064, and 3428 Å.) were well developed.

*Flame Spectra at Higher Pressures*—Our spectrograms for flames maintained at 10, 30, and 100 atmospheres respectively are reproduced in figs 4-6, Plates 7 to 9. The principal effects thereon (i) of increasing pressure with each burning medium, and (ii) of successive CO by H<sub>2</sub> replacements at each pressure, are summarized in Table I.

Reviewing the results as a whole we find —

*Effects of Pressure* (1) *With a Pure CO-Flame*—While the continuous spectrum was (if anything) intensified and completely obliterated the CO-O<sub>2</sub> bands as the pressure was increased up to 100 atmospheres, the 3064 HO-bands visible at atmospheric pressure became very faint at 10, and completely faded out at 30 atmospheres and above, indicative of the CO-oxidation becoming increasingly direct with rising pressure.

(2) *With a Pure H<sub>2</sub>-Flame* the general effect of increasing pressure was to strengthen all the HO-bands, and at 100 atmospheres to produce reversals at the head of the 3064 Å band—doubtless owing to the survival of free HO-radicals in the cooler atmosphere surrounding the flame. And at the same time the bands did not degrade so rapidly, indicative of a progressive shifting of the energy distributed away from the band-head as the pressure rose. Also there was a growing tendency, most marked at 100 atmospheres, to produce a slightly diffused background. There was no sign of any H-lines developing.

(3) *With CO-H<sub>2</sub>-Flames*—With corresponding mixture-compositions increasing pressure induced a growing tendency for the HO-bands to become relatively fainter and for the continuous CO-spectrum to become more pronounced, as though the direct oxidation or carbonic oxide was progressively superseding the indirect. This is well shown in fig. 7, Plate 10 in which the spectra for a 78CO/22H<sub>2</sub> flame at 1, 10, 30, and 100 atmospheres respectively are reproduced. In other words, the effect, first observed by Weston, of successive H<sub>2</sub>-additions to a CO-flame at atmospheric pressure are now seen to be counteracted by increasing pressure.

Combustible  
gas  
 $\left\{ \begin{array}{l} \text{CO} \\ \text{H}_2 \end{array} \right.$   
100 0

93 7

78 22

50 50

22 78

0 100

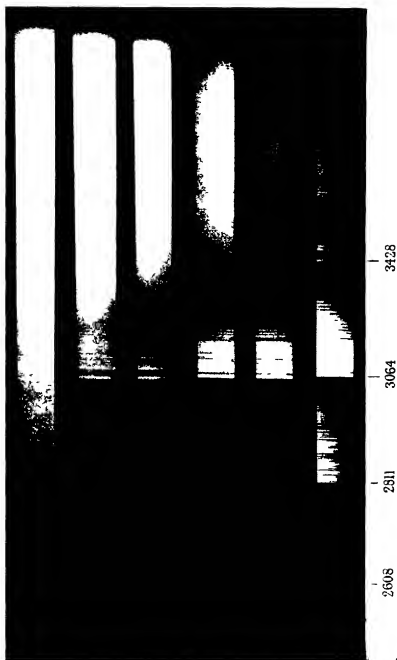


FIG 3—Spectra of carbonic oxide-hydrogen flames burning in oxygen at atmospheric pressure

Combustible  
gas

$\left\{ \begin{array}{l} \text{CO} \quad \text{H}_2 \\ 100 \quad 0 \end{array} \right.$

93 7

78 22

50 50

22 78

7 93

0 100

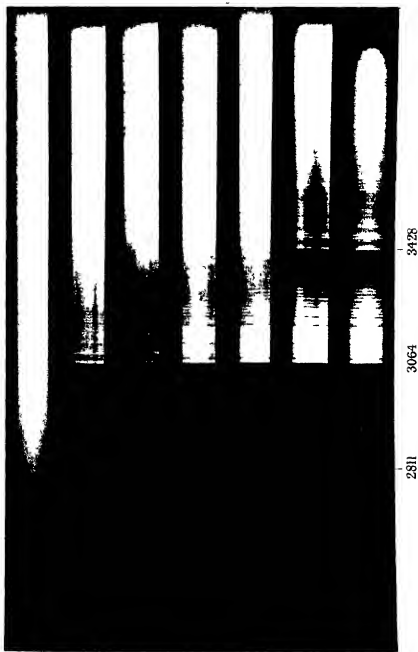


FIG. 4.—Spectra of carbon monoxide flames burning in oxygen at 10 atmospheres pressure

Combustible  
gas

CO H<sub>2</sub>  
100 0

93 7

78 22

50 50

22 78

7 93

0 100

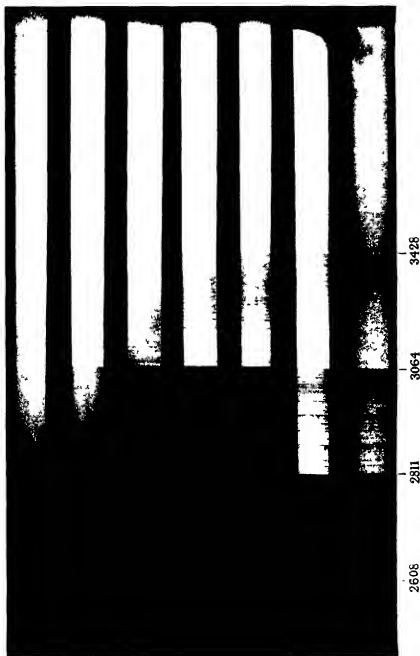


FIG. 5.—Spectra of carbonic oxide-hydrogen flames burning in oxygen at 30 atmospheres pressure.

Combustible  
gas  
 $\left\{ \begin{array}{l} \text{CO} \quad \text{H}_2 \quad \text{O} \\ 100 \quad 0 \end{array} \right.$

93 7

78 22

50 50

22 78

7 93

0 100



2811

3064

3458

FIG 6 — Spectra of carbon monoxide flames burning in oxygen at 100 atmospheres pressure



Fig 7-78 CO-H<sub>2</sub> mixture





Table I—Spectra of CO-H<sub>2</sub>-O<sub>2</sub> Flames OH bands observed 0.9 at 3064 Å., 1.0 at 2811 Å., 2.0 at 2608 Å., and 0.1 at 3438 Å.

Fuel		Pressure			Effect of pressure on given mixture	
CO %	H <sub>2</sub> %	Atmospheric	10 atmospheres	20 atmospheres		100 atmospheres
100	0	Strong continuous spectrum extending to far on 2400 Å. 0.0 band variable	Strong continuous spectrum extending to far on 2400 Å. 0.0 band faintly variable	Strong continuous spectrum, no OH bands visible	Strong continuous spectrum, no OH bands visible	Little change in continuous spectrum, 0.0 band fades out at 20 atmospheres.
93	7	Slight reduction in continuous spectrum 0.9 and 1.0 bands variable	Strong continuous spectrum, 0.0 band faint 1.0 band faint	Strong continuous spectrum, 0.0 and 1.0 bands variable 1.0 band very faint	Strong continuous spectrum, 0.0 band faint 1.0 band faint 1.0 band reversed at head	Reduction of continuous spectrum by addition of H <sub>2</sub> at 1 atmosphere (Counteracted by pressure
76	22	Much less continuous spectrum, 0.0 and 1.0 bands very distinct	Reduced continuous spectrum, 0.0 and 1.0 bands weakly visible	Slightly reduced continuous spectrum, 0.0 band strong, 1.0 band weak	Strong continuous spectrum, 0.0 band faint 1.0 band reversed at head	Increased continuous spectrum with pressure, 1.0 band fades out above 50 atmospheres
50	50	Continuous spectrum very much reduced 0.0 and 1.0 bands very strong, 2.0 and 0.1 bands now visible	Reduced continuous spectrum, 0.0 and 1.0 bands strong, 2.0 and 0.1 bands weak	Reduced continuous spectrum, 0.0 and 1.0 bands strong, 2.0 band weak	Strong continuous spectrum, 0.0 band strong, 1.0 band faintly variable	At 1 atmosphere spectrum resembles that of H <sub>2</sub> flame, at 100 atmospheres that of CO flame
22	78	Very little continuous spectrum, 0.0, 1.0, 2.0 bands strong, 0.1 band weak	More continuous spectrum than at 1 atmosphere, 0.0 and 1.0 bands strong, 2.0 and 0.1 bands weaker	Reduced continuous spectrum, 0.0 and 1.0 bands strong, 2.0 and 0.1 bands weak	Much continuous spectrum, 0.0 band reversed at head, 1.0 and 0.1 bands variable	Decided pressure effect in increasing continuous spectrum and reducing intensity of bands
7	93	---	Little continuous spectrum, 0.0, 1.0, and 0.1 bands strong, 2.0 band weak	Some continuous spectrum, 0.0, 1.0, 2.0, and 0.1 bands all very distinct	Bands very heavy	Continuous spectrum increased with pressure, intensity of bands reduced
0	100	No continuous spectrum, 0.0, 1.0, 2.0, and 0.1 bands all very clear	No continuous spectrum, bands very strong and do not degrade so rapidly	Bands very heavy	0.0 band reversed at head, slight diffuse background	At higher pressure bands do not degrade so rapidly
Effect of composition combustible.		Bone-Watson effect well marked.	Bone-Watson effect not so well marked	Bone-Watson effect no longer evident	Continuous spectrum very strong even with large proportions of H <sub>2</sub> in the fuel.	---

Speaking generally, although in some of the spectrograms of  $\text{CO-H}_2$  flames at atmospheric pressure the banded spectrum of burning carbonic oxide underlying its continuous spectrum is faintly discernible in the original photographic plates, in those obtained at higher pressures it is swamped by the intensity of the continuous spectrum

In conclusion, we desire to acknowledge our indebtedness (i) to Dr D M Newitt for help in connection with the design of the high pressure combustion chamber employed in these experiments, and (ii) to the Government of Northern Ireland for a grant which has enabled one of us (F G L.) to devote his whole time to the work

### *Summary*

A new apparatus is described for maintaining continuous flames of combustible gases in air or oxygen at high pressures, and with it has been studied the influence of increasing pressure up to 100 atmospheres upon the flame-spectra of hydrogen and carbonic oxide, and various mixtures thereof, burning in oxygen

(1) *With a Pure CO-flame*—While the continuous spectrum was (if anything) intensified and completely obliterated the  $\text{CO O}_2$  bands as the pressure was increased up to 100 atmospheres, the 3064 HO-bands visible at atmospheric pressure became very faint at 10, and completely faded out at 30 atmospheres and above, indicative of the  $\text{CO}$ -oxidation becoming increasingly direct with rising pressure

(2) *With a Pure  $\text{H}_2$ -flame*—The general effect of increasing pressure was to strengthen all the HO-bands, and at 100 atmospheres to produce reversals at the head of the 3064 Å band—doubtless owing to the survival of free HO-radicals in the cooler atmosphere surrounding the flame. And at the same time the bands did not degrade so rapidly, indicative of a progressive shifting of the energy distributed away from the band-head as the pressure rose. Also there was a growing tendency most marked at 100 atmospheres to produce a slightly diffused background. There was no sign of any H-lines developing

(3) *With  $\text{CO-H}_2$ -Flames*—With corresponding mixture-compositions increasing pressure induced a growing tendency for the HO-bands to become relatively fainter and for the continuous  $\text{CO}$ -spectrum to become more pronounced, as though direct oxidation of carbonic oxide was progressively superseding the indirect. In other words, the effect, first observed by Weston, of successive  $\text{H}_2$ -additions to a  $\text{CO}$ -flame at atmospheric pressure are now seen to be counteracted by increasing pressure

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## *The Supposed Intervention of Steam in Hydrocarbon Combustion*

By WILLIAM A BONE, F R S, and JACK BELL, B Sc, A R C S

(Received November 15, 1933)

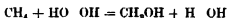
[PLATES 11-12]

An important issue in connection with the theory of hydrocarbon combustion, needing further discussion, is whether or not the presence of water vapour plays any essential role in the process.

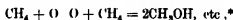
H E Armstrong, who originated the hydroxylation theory in 1874, has always insisted that it does, its action (taking the first oxidation stage of methane as a typical example) being supposed as follows —



the hydrogen peroxide so formed participating in the oxidation and regenerating the steam thus --



On the other hand, Bone and his co-workers, while maintaining that the oxidation involves successive "hydroxylation" stages, regard the action of the oxygen (whether molecular or atomic) as direct, thus --



or possibly, in flames,  $\text{CH}_4 + \text{O} = \text{CH}_3\text{OH}$ , etc

As H E Armstrong has recently re-affirmed his view in discussing Bone's experimental results, the time seems opportune for both reviewing the evidence already published and supplementing it by some newly acquired in our laboratories

### *1 Previous Evidence about Slow Combustion*

Previous evidence all relates to non-explosive slow combustion and comprises two somewhat different sets of conditions, which had best be dealt with in the reverse of their chronological order, as follows (A) in the course of recent re-investigations (1930-33) in our laboratories of the slow combustions

\* This equation expresses two outstanding facts, namely, (i) that the slow combustion  $2\text{CH}_4 + \text{O}_2$  is much the most reactive of all methane oxygen mixtures, and (ii) that methyl alcohol is the first recognisable product

of methane, ethane, and ethylene,\* studies were made of the effects of small additions of water-vapour upon the durations of both "induction" and "reaction" periods of selected  $P_2O_5$ -dried hydrocarbon-oxygen mixtures in a quartz vessel at atmospheric pressure and a temperature no higher than was required to give a conveniently measurable rate of reaction in each case. The reaction vessel had been thoroughly dried-out by prolonged evacuation at the experimental temperature prior to the admission of the gaseous medium, which was slowly passed in through a metre long column of pure redistilled phosphoric anhydride. The drying thus effected, while not severely intensive, would be as thorough as was needed for the purpose in view. In the "moist" experiments, 1 or 2% by volume of vapour from "ammonia-free" redistilled water was added to the so-dried medium just as it entered the reaction vessel. The observed results are given in Table I.

Table I

Mixture employed	Temperature of reaction vessel	Hygroscopic condition	Observed durations of-	
			'Induction' (mins.)	'Reaction' (mins.)
$2C_2H_4 + O_2$	447	$\left\{ \begin{array}{l} \text{Dry} \\ \text{Moist, 2\%} \end{array} \right.$	$\left\{ \begin{array}{l} 6 \text{ to } 7 \\ 1 \text{ to } 4 \end{array} \right.$	$\left\{ \begin{array}{l} 91 \text{ to } 98 \\ 15 \text{ to } 36 \end{array} \right.$
$C_2H_6 + O_2$	316	$\left\{ \begin{array}{l} \text{Dry} \\ \text{Moist, 1\%} \end{array} \right.$	$\left\{ \begin{array}{l} 30 \\ 10 \end{array} \right.$	$\left\{ \begin{array}{l} 70 \\ 25 \end{array} \right.$
$C_2H_4 + O_2$	300	$\left\{ \begin{array}{l} \text{Dry} \\ \text{Moist, 1\%} \end{array} \right.$	$\left\{ \begin{array}{l} 54 \\ 50 \text{ to } 60 \end{array} \right.$	$\left\{ \begin{array}{l} 50 \text{ to } 52 \\ 50 \end{array} \right.$

It is thus seen that while additions of water-vapour to the  $P_2O_5$ -dried  $2C_2H_4 + O_2$  and  $C_2H_6 + O_2$  (*i.e.*, the paraffin-containing) media materially shortened both the induction and reaction periods, they had little or no effect upon the  $P_2O_5$ -dried  $C_2H_4 + O_2$  (*i.e.*, olefine-containing) medium. None of the dried media, however, appeared unreactive.

(B) In 1906 Bone and Andrew published† the results of a careful study of effects of an intensive  $P_2O_5$ -drying (4 to 12 weeks) upon the reactivities of  $C_2H_6 + O_2$ ,  $C_2H_4 + O_2$ , and  $C_2H_2 + O_2$  media at various temperatures between  $300^\circ$  and  $500^\circ$  and pressures between 2 and 3 atmospheres. For details of

\* 'Proc. Roy. Soc.,' A, vol. 129, p. 434 (1930) vol. 134, p. 578 (1932)

† 'J. Chem. Soc.,' vol. 80, pp. 652-659 (1906)

the experimental procedure, the original paper should be consulted, suffice it now to say that on comparing the behaviour of each *dried* and *undried* medium, when subjected to the same degree and duration of heat, little or no difference could be found between them—the *dry* always reacting quite as fast as (and if anything even slightly faster than) the corresponding undried medium. This evidence was strengthened by the fact that the same drying procedure had caused enormous differences between the percentages of reaction (e.g., nil and 42, 0.5 and 52, 3 and 40 or 45) in “dry” and “undried”  $2\text{H}_2 + \text{O}_2$  media, respectively, after 15 minutes at  $525^\circ$ . This is well illustrated by the results of two such comparative sets of experiments with a  $\text{C}_2\text{H}_4 + \text{O}_2$  medium, where the “drying period” extended over 11 weeks. The temperature at which the first visible sign of reaction occurred, when the tubes were subsequently heated, was about the same ( $385^\circ$ – $398^\circ$  C) in the “dry” as in the “moist” medium, and after 15 minutes further heating up to a maximum of  $430^\circ$  to  $450^\circ$ , the amounts of reaction were substantially the same in each, as can be seen from Table II.

Table II—Results with “Moist” and “Dry”  $\text{C}_2\text{H}_4 + \text{O}_2$  Media at  $430^\circ$  to  $450^\circ$  and 2.5 Atmospheres for 15 minutes (Bone and Andrew)

State of medium	Percentage contraction on cooling	Percentage composition of gaseous products					
		$\text{CO}_2$	$\text{CO}$	$\text{C}_2\text{H}_4$	$\text{C}_2\text{H}_6$	$\text{H}_2$	$\text{O}_2$
(1) $\text{P}_2\text{O}_5$ -dried, 11 weeks	37.0	10.25	45.75	32.65	2.10	1.05	8.20
	37.7	10.25	44.50	28.85	6.35	4.00	6.05
(2) Moist, $\text{H}_2\text{O} = 2\%$	34.8	10.60	48.4	27.6	4.65	2.70	6.05
	36.1	11.45	50.8	24.85	5.15	3.25	4.50

Two experiments with a 4-weeks  $\text{P}_2\text{O}_5$ -dried  $\text{C}_2\text{H}_4 + \text{O}_2$  medium showed in (1) first visible signs of reaction after 27 minutes at  $315^\circ$ , and complete disappearance of oxygen within a further 8 minutes during which the temperature had risen to  $415^\circ$  ( $p_2/p_1 = 0.672$ , and gaseous products contained  $\text{CO}_2 = 19.65$ ,  $\text{CO} = 56.65$ ,  $\text{C}_2\text{H}_4 = 22.5$ ,  $\text{C}_2\text{H}_6 = 0.35$ ,  $\text{H}_2 = 0.85$ , and  $\text{O}_2 = \text{nil } \%$ ), while in (2) when the contents of the tube were heated to  $520^\circ$ , they exploded after 2 minutes with shattering effect.

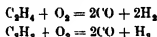
Two sets of comparative experiments with “dry” and “undried”  $\text{C}_2\text{H}_4 + \text{O}_2$  mixtures, respectively, at temperatures between  $415^\circ$  and  $450^\circ$  left no room for doubt but that reactivity was definitely increased by a 7 weeks

**P<sub>2</sub>O<sub>5</sub>-drying** Ignition finally occurred in both "dry" media, and also in one of the two moist ones

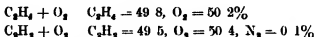
Although the experimental conditions precluded any estimation of the effects of the intensive drying on the length of the "induction period" in each case—which would in any case be extremely short—there could be no doubt about the P<sub>2</sub>O<sub>5</sub>-drying having (if anything) rather increased than diminished the reactivities of all the media examined at the temperatures and pressures concerned

### II *New Evidence re Explosive Combustion*

Although the evidence regarding slow combustion seemed decisive enough on the point at issue, it was thought desirable to extend the enquiry to explosive combustion, and for this purpose equimolecular mixtures of ethylene or acetylene and oxygen seemed most suitable\* inasmuch as on explosion they yield substantially nothing but carbonic oxide and hydrogen, without any discernible carbon separation or steam formation, thus —



*The Experimental Mixtures*—The experimental mixtures were made up very accurately over dry redistilled mercury in a large cylindrical gas burette A, fig 2, from stocks of highly purified hydrocarbon and oxygen, each hydrocarbon having previously undergone (i) a thorough chemical purification followed by (ii) two successive liquefactions and fractional vapourizations of the liquid, the middle third only being reserved in each case, on analysis the compositions of the experimental mixtures were found to be as follows —



*The Explosion Tubes*, GG, fig 1, were constructed out of Jena red-line borosilicate glass tubing (internal diameter = circa 2 cm), each was 35 cm long, having expanded ends for the reception of the drying-agent, and being fitted mid-way with "swinging" platinum electrodes (1 mm diameter terminating in a bulb 1.5 mm diameter)

*Cleaning and Drying of the Explosion Tubes*—The elaborate ritual for thoroughly cleaning and drying out the innares of the explosion tubes was that

\* Experiments on CH<sub>4</sub> + O<sub>2</sub> or C<sub>2</sub>H<sub>6</sub> + O<sub>2</sub> mixture were deemed unsuitable because they would have yielded H<sub>2</sub>O on explosion.

followed in previous similar experiments in our laboratories and has been fully described in a former publication. Between the initial and final stages, and again thereafter, the platinum electrodes were electrically "glowed-out" each time *in vacuo* for 8 hours and then in 2 mm of oxygen for 3 hours, for the elimination of any trace of occluded hydrogen. All other glass apparatus used during the experiments was similarly cleaned and dried.

*Final Drying of the Experimental Mixtures*—Finally the clean and dried explosion tubes, GG, into each expanded end of which a quantity of purified and redistilled  $P_2O_5$  had been introduced, were connected through a suitable KOH- and  $P_2O_5$ -drying train, CDE, and manometer, F, with the gas burette

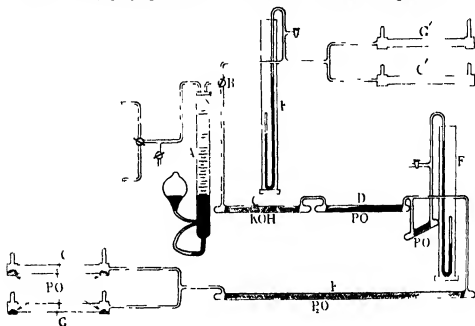


FIG 1

A containing the experimental mixture, as shown in fig 1. An alternative connection was also provided, through the tap B, between the gas burette and the explosion tubes G' G' used in the comparative experiments with the corresponding "moist" mixtures, which were about half-saturated with water vapour at the room temperature ( $17^\circ$  to  $20^\circ$ ). The actual drying system comprised (i) a 50-cm long tube packed with solid KOH followed in succession by two tubes 50 and 100 cm long respectively, filled with purified and redistilled  $P_2O_5$ . All joints throughout the apparatus were of fused glass, the fusion being effected in a dust free and  $CaCl_2$ -dried  $CO-O_2$  blowpipe flame.

Before transferring the gaseous mixture from the gas burette A to the explosion tubes, GG, the whole of the intervening apparatus was exhausted down to a pressure of 0.001 mm by means of a Hyvac pump which was kept continually running each time for at least 24 hours. Finally the mixture was passed, at a rate not exceeding 1 and sometimes as low as 0.5 c.c. per minute, from the gas burette, through the "drying system" into the explosion vessels. The filling arrangements in the control experiments with the corresponding "undried" mixture were similar to the foregoing, except for the omission of the "drying train," and of  $P_2O_5$  in the explosion tubes concerned.

On completion of the filling at room temperature and the selected firing pressure (735 mm. and  $20^\circ$  for the  $C_2H_4 + O_2$  and 370 mm. and  $17^\circ$  for the  $C_2H_2 + O_2$  mixture), the tubes were finally sealed off in the blow-pipe at a constriction previously made in the glass for the purpose.

*The Drying Period*.—The subsequent "drying period" always extended over rather more than 300 days during which the expanded ends of each tube were frequently tapped to expose a fresh  $P_2O_5$  surface to the gaseous medium, and at regular intervals of 14 days the main part of the tube between the expanded ends (but not the latter) was heated to about  $150^\circ$  in order to ensure the dispersion of any film of moisture that otherwise might have adhered to the inner walls.

*Control Experiments*.—While the drying was in progress, two special control experiments were made as a precaution to ensure that prolonged exposure to the  $P_2O_5$ -drying agent employed does not alter the composition of either a  $C_2H_4 + O_2$  or a  $C_2H_2 + O_2$  mixture—by some preferential absorption of one or other of its constituents. In neither case, however, had a 7 months' exposure any appreciable effect upon the composition of the mixture.

#### *Photographic Investigations of the Explosions*

Two pairs of "dry" and "moist" tubes (the latter for comparison) for each of the two experimental mixtures having been so prepared, and the drying period ended, arrangements were made for firing each pair of tubes in a horizontal position by means of a condenser discharge of 0.06 m.f. at 1000 volts and photographing the resulting explosion on a film moving vertically, at constant velocity, in one of our Fraser high-speed cameras. In this way comparative photographic analyses were obtained of the flame movements throughout each pair of "dry" and "moist" explosions, from which a series of comparative flame-speeds could be calculated, showing precisely what had



been the effect (if any) of the intensive  $P_2O_5$ -drying. It will, however, suffice to reproduce and analyse the relative photographs for *one* pair only of "dry" and "moist" explosions for each of the two experimental mixtures, and then merely to tabulate, as confirmatory evidence, the relative flame speeds observed in the other pair of explosions with each mixture.

A *The  $C_2H_4 + O_2$  Explosions*—(1) The relative photographs for the first pair of "dry" and "moist" explosions (tube internal diameter = 2.0 cm) are reproduced in figs. 2 and 3, Plate 11.

The film speeds were 33.0 for the "dry" and 31.8 metres per second for the "moist" explosion, respectively. The drying period in the former had been 316 days, and the gaseous pressure in both cases 736 mm. On firing the mixtures, in each an explosion ensued instantaneously, the flame traversing the mixture much faster than the eye could follow and ultimately filling the whole tube, and while the tube itself always remained intact, its capillary end was fractured by the explosion.

The resulting photograph showed that the flame speed had been materially faster and the total duration of luminosity (T.D.) also shorter in the "dry" than in the "moist" explosion. Each photograph shows clearly (i) the flame starting instantaneously from the igniting discharge and thence spreading rapidly and symmetrically towards each end of the tube, (ii) a continuance of luminosity for some time after the flame had completely filled the tube, during which the medium was traversed from end to end by compression waves, and (iii) the production in the central region round the electrodes and also in the flame front itself, of striæ of enhanced luminosity, all of which coalesced just as the advancing flame fronts reached the ends of the tube.

Particulars of the flame-speeds deduced from the photographs are as follows:—

Average flame speed from source to point	(a) "Dry" explosion (metres per sec.)	(b) "Moist" explosion (metres per sec.)
(i) 5 cm. distant	55.6	54.4
(ii) 10 cm. distant	68.8	70.4
(iii) To end of tube	70.0	55.0
Total duration of luminosity	(milliseconds) 6.3	(milliseconds) 7.7

(2) The internal diameter of the tubes used in this pair of explosions was 2.3 cm., otherwise the conditions were precisely the same as previously.

Particulars of the flame-speeds deduced from the resulting photographs are as follows —

Average flame-speed from ignition to point	(a) "Dry" explosion (metres per sec)	(b) "Moist" explosion (metres per sec)
(i) 5 cm distant	65.0	53.8
(ii) 10 cm distant	89.4	61.7
(iii) End of tube	85.5	60.0

It is thus seen that in each pair of explosions the average flame speed through the "dry" was decidedly (27%–42%) greater than through the "moist" medium.

**B The  $C_2H_2 + O_2$  Explosion**—(1) The relative photographs for the first pair of "dry" and "moist" explosions (internal diameter of tube = 2.0 cm) are reproduced in figs 4, 5, Plate 12. The drying period in the former had been 314 days and the pressure in both cases 371 mm.

In each case a deafening report immediately followed the passage of the discharge, the tube being shattered by the force of the explosion, although it had remained intact until completely filled with flame.

The film speeds were 79.7 for the "dry" and 74.0 metres per second for the "moist" explosion respectively. The average flame speeds were very high, namely, 700 and 585 metres per second respectively, or about 20% higher in the "dry" than in the "moist" medium. The fact that the total duration of luminosity was apparently greater in the "dry" than in the "wet" explosion is probably of no particular significance owing to the fracturing of the tube after the flame had reached its ends.

Particulars of the flame speeds deduced from the photographs are —

Average flame-speed from ignition to point	(a) "Dry" explosion (metres per sec)	(b) "Moist" explosion (metres per sec)
(i) 5 cm distant	416	333
(ii) 10 cm distant	476	417
(iii) End of tube	700	585

(2) The internal diameter of the tubes used in the second pair of explosions was 1.8 cm, but otherwise the conditions were the same as previously. Particulars of the flame-speeds deduced from the resulting photographs are given in the following table.

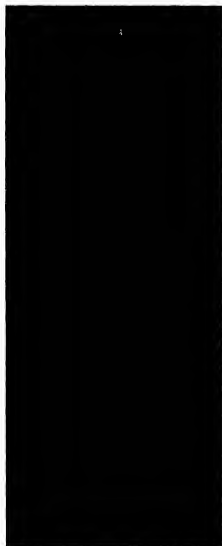


FIG 2 — "Dry "



FIG 3 — "Moist "

$C_2H_4 + O_2$  explosions

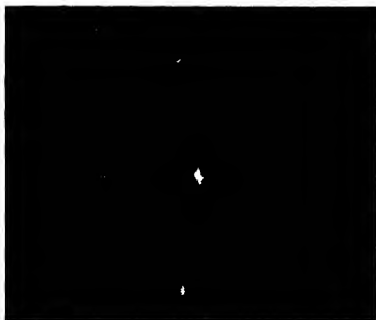


FIG 3 — Moist



$C_2H_2 - O_2$  explosions

FIG 4 — Dry

Average flame-speed from ignition to point.	(a) "Dry" explosion (metres per sec.).	(b) "Moist" explosion (metres per sec.)
(i) 5 cm. distant	312	278
(ii) 10 cm. distant	434	385
(iii) End of tube	600	500

It is thus seen that in both cases the flame-speeds were greater in the "dry" than in the "moist" explosions. Another point of interest about the photographs to which attention may be directed is the entire absence of any sign of carbon separation in the flame front, which accords well with the view that the primary reaction is  $C_2H_2 + O_2 \rightarrow (C_2H_2O_2) \rightarrow 2CO + H_2$ .

### Conclusion.

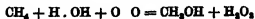
(1) The results as a whole are, we think, decisive against any supposition that in general the combustion of a hydrocarbon is conditioned by the presence and chemical intervention of steam, as H. E. Armstrong supposes, although in certain but not every case of slow combustion it may in some way assist the reaction. Hence we conclude that the oxidation is, in general, direct.

(2) Nor yet do photographs of the  $C_2H_4 + O_2$  and  $C_2H_2 + O_2$  explosions lend any support to the view (sometimes suggested) that the final production of substantially carbonic oxide and hydrogen only is nearly the resultant of the successive reactions  $C_2H_4 + O_2 \rightarrow 2C + 2H_2O \rightarrow 2CO + 2H_2$ , and  $C_2H_2 + O_2 \rightarrow CO + C + H_2O \rightarrow 2CO + H_2$ , for they show neither carbon separation in the flame front, nor yet any sign of carbon-steam reaction behind it.

We desire to thank the Trustees of the Walker Scholarship Fund, the Miners Welfare National Scholarship Scheme, and the D. S. I. R. for grants which have enabled one of us (J. B.) to participate in the research.

### Summary

The paper deals with the question as to whether in hydrocarbon-combustion the oxygen acts directly in hydroxylating the hydrocarbon or only indirectly (as H. E. Armstrong maintains) through the intervention of steam, as, for example —



Existing evidence in regard to the effects of  $P_2O_5$ -drying upon the slow combustion of typical hydrocarbon-oxygen mixtures is reviewed, and new evidence in regard to its effects upon explosions of  $C_2H_2 + O_2$  and  $C_2H_4 + O_2$  media is detailed. The conclusion is reached that intensive  $P_2O_5$ -drying if anything increases rather than diminishes the reactivities of hydrocarbon-oxygen mixtures, both in slow and explosive combustion, and therefore that steam does not intervene, the oxidation being direct

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### *The Bending of Marble*

By Lord RAYLEIGH, For Sec. R S

(Received December 1, 1933)

[PLATE 13]

#### § 1 *Introduction*

Cut slabs of marble are often noticeably bent. An old marble mantelpiece, say 1.5 metres long, will always show a depression in the middle when tested with a stretched thread. This depression is often 1 to 2 mm, and I have heard of cases where it is much more. My interest in this phenomenon was aroused by a conversation with the late Dr D. W. Dye, F.R.S., and a few preliminary experiments were made. My first notion was that the marble bent by viscous flow, as a stick of sealing wax will do when loaded in the middle, and I even speculated as to whether it would be possible by the continued application of force to bend a piece double. The well-known experiments of Adams and Nicholson\* on the flow of marble under high pressures seemed at first glance to give this notion some plausibility.

Bent pieces of marble were obtained from a local marble merchant, Mr J. B. Slythe, of Witham. He was not able to say anything definite about their origin, except that they had been in his yard for many years. A bent strip was supported at the ends with the concave side up, and loaded at the middle with as much weight as it was judged safe to put on. A suitable index

\* 'Phil. Trans.,' vol. 195, p. 363 (1901).

was fixed at the middle and observed by a microscope with eyepiece scale. There was noticeable motion the first day, but it rapidly diminished in amount from day to day, until it practically ceased after a few weeks. I then reversed it, thinking that if there were a limit to the bending it would at least be possible to restore its original straightness. Again, there was noticeable motion at first, but it soon almost stopped, and calculation showed that decades would be required at the least to straighten the piece in this way, even under the favourable conditions of the experiment. This made it improbable that the piece could have been bent cold. It then seemed that it might have been bent by the application of force while hot. A similar experiment was accordingly tried at a higher temperature. A strip of new marble was placed on the hot-water pipes, similarly supported at the ends and heavily loaded in the middle. It was covered with cotton-wool lagging, and thus maintained at from 45° to 50° C throughout the winter. But tests with a spherometer showed that only very slight bending had occurred, and this was nearly all in the first week or two.

At this stage it was difficult to know what to think, the more so that Mr Slythe informed me that he had had pieces of marble which would gradually bend under their own weight when leaned against the wall, and would become straight again in course of time, if reversed. I eventually succeeded in obtaining a slab, obviously an old mantelshelf, of very marked curvature, which showed this behaviour. Careful observations were made on it. The length was 182.4 cm, the breadth 15 cm, and the thickness 2 cm. It was geometrically supported at the ends, and loaded at the middle with weights amounting to 8.6 kg, and a suitable pointer and mirror glass scale provided. The weights were removed for taking readings. The first tests were made with the smooth side up, i.e., as used for a mantelshelf. In seven weeks the index had sunk 6.1 mm, increasing the *sagitta*\* of the arc from 16.9 mm to 23 mm. Towards the end of this period the motion had become very slow, though it could still be detected in a week.

Having got something near the maximum curvature in this way, the slab was reversed and replaced with the convex (rough) side upwards, in order to straighten it. After reversal, when the slab had been allowed to take up its position, most of the concavity of the smooth side had disappeared. At first reading the *sagitta* was only 8.7 mm. This great diminution of curvature had taken place immediately under the weight of the slab acting in the reversed

\* This convenient word for the maximum distance from the chord to the arc is used in Newton's "Principia," and is here revived.

sense Leaving the slab under its own weight the sagitta was further diminished to 6.7 mm. in 5 days The change had become very slow, and the same weights as before (8.7 kg.) were put on This resulted in a further change, and eventually the slab, as observed unloaded, became straight, and finally achieved a curvature in the opposite direction (sagitta—2.5 mm.) The motion under the load had then become very slow, amounting to 1.7 mm. in 77 days, and the experiment was discontinued

It is noticeable that this specimen, unlike the previous one, is highly flexible within certain limits, and that these limits are unsymmetrical It can take a much greater concavity on the smooth side (i.e., the top side of mantelshelf) than on the rough side This may be regarded as a combination of permanent curvature with flexibility

Finally, a marble strip was brought to me which had been in a wooden shed burnt to the ground in an accidental fire, and which had presumably become considerably heated This had very marked flexibility without permanent curvature

It appears therefore that flexibility and permanent curvature (warping) may appear either combined or separately, and that at all events in some striking cases the specimens which show either have been subjected to a more or less high temperature\* It was with this clue that some success was obtained in the further investigation

Preliminary experiments on baking marble in a kitchen oven at about 300° C. showed that marked flexibility resulted from this treatment A demonstration experiment was shown at the Royal Society soirée on June 21, 1933 The strips exhibited were each gripped at one end in a wooden vice, and protruded some 30 cm The thickness was about 8 cm. The baked strip could be moved to and fro through a range of 2 cm. between stops, by gentle pressure with the forefinger The unbaked strip provided for comparison was rigid

## § 2 *Systematic Observations on Flexure and Expansion, after Baking at Various Temperatures*

For a closer examination apparatus was arranged as in fig. 1 The marble was vertical. It was gripped at its upper end in a cast iron vice of special construction, carried on the wall The lower end could be pulled to right or left by the attached threads By this method any complication from the

\* Mr. Slythe was inclined to think that old marble, i.e., marble which had been long exposed to the weather, as tombstones and the like, often showed flexibility I have not so far been able to confirm this.



weight of the marble itself is avoided, and the force admits of easy reversal. The threads passed over pulleys working on centres with a minimum of friction, and carried the scale pans. Weight in one or other of these pans provided a known force. A brass piece carrying two separate index needles

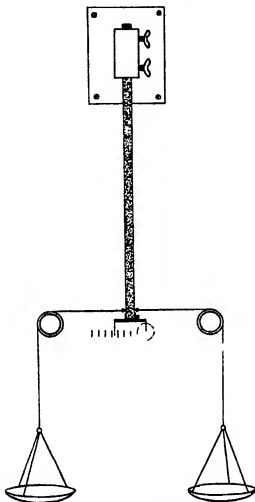


FIG 1 ( $\frac{1}{2}$  actual size)

was clamped to the lower end of the marble quite independently of the attachment for the threads, which latter is not shown in detail. One of the needles is the index for minute movement, and is read by a microscope with eyepiece scale [Microscope shown by the dotted circle]. The other needle is read by

a millimetre scale on mirror glass, and is used for indicating the larger movements. The use of two separate indexes is convenient since the microscope need not be removed when the millimetre scale is used. The microscope is useful in the early stages of the experiment to examine the small flexure of unbaked marble, or marble baked at low temperatures.

Any shift of the marble in the vice would make the results inaccurate for the smaller deflections. To avoid crushing the marble, strawboard packings 2.5 mm thick were used on either side between it and the iron jaws, which latter were screwed up "thumb tight" by means of the three thumb screws. [The details are shown in fig. 2. The lower thumb screw shown is to be interpreted as representing two, one above, one below, the plane of the diagram.]

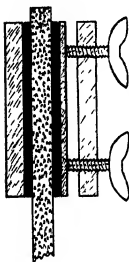


FIG. 2 ( $\frac{1}{2}$  actual size)

As a test of the reliability of holding, a piece of flat bar iron of about the same dimensions was set up instead of the marble, and clamped in the same way. After loading with 1 kilogram, and unloading again, the original microscope reading was exactly recovered. It is concluded from this that the method of holding is satisfactory, and that the strawboard packings are unobjectionable.

For baking the marble strips an electrical oven was constructed. An asbestos muffle received the strip. This muffle was wound with resistance wire, and surrounded with a thick layer of slag wool for heat insulation. The furnace current was increased a definite amount for each experiment and the specimen heated for about 20 hours, the temperature being taken at the end of that time. Some

hours are required to reach a steady temperature, so that the baking was not under strictly uniform conditions.\*

As is known, marble undergoes a permanent increase of length on heating. It was considered desirable for reasons which will appear later to observe the increment concurrently with the mechanical tests. For this purpose fine scratches were made at the two ends of the marble strip, about 400 mm apart. The marble was laid on a planed iron slab, and a piece of sheet glass laid on the marble. The glass had fiducial lines engraved upon it about 399 mm

\* It is possible that alternate heating to a given temperature and cooling might have more effect than continuous heating to the same temperature. This point has not been examined.

apart, thus 1 mm. less than the distance between the scratches on the marble. The engraved side of the glass was, of course, downwards, and weights were placed on it to keep the whole flat and steady. The short distance at each end between the mark on the glass and that on the marble was measured with travelling microscopes. In this way the small increments of length were determined. The greater part of the length was taken care of by the glass standard, its exact absolute value being unimportant.

A strip of marble, believed to be from Carrara, was taken, measuring 78 mm broad, 9 mm thick, and a little over 400 mm long. It was clamped in the vice, the protruding length being 34 cm and was carried through a cycle of changes, being first loaded up to +1000 grams (i.e., 1000 grams on the right-hand pan). This was taken off 200 grams at a time, with readings at each step. Weights were then progressively added on the other side to -1000 grams, and so on until two complete cycles had been gone through, the mean of the two cycles being taken. The discrepancies were small, particularly in the early stages of the experiment. Having gone through this process with the marble in its initial condition, it was baked at a determined temperature, on cooling the increase of length was measured, and the mechanical tests repeated, and so on, increasing the baking temperature each time, until finally the specimen broke off under the standard force of 1 kilogram.

The readings were always taken as quickly as convenient after loading or unloading. There is a certain creep following on this, which goes on for a long time, becoming slower and slower, and perhaps never definitely ceasing. But the initial movement is the important one, the creep being comparatively small. In the cyclic tests the creep was ignored, though it probably accounts for any small outstanding lack of definiteness in the measurements.\*

The most important part of the results is in the following table.

The total flexure referred to is the difference between the extreme positions under load. The inelastic flexure is the difference between the positions on unloading after pulling to the right and left respectively.

The small permanent elongation observed after baking at 55° is doubtful. Definite elongation is first observable after baking at 77°, and at this point the change of mechanical properties also begins to appear.

It will be noticed that the inelastic flexure becomes a much larger fraction of the whole as the baking temperature is increased. With the load used ( $\pm 1$

\* This creep was the subject of observation in my earlier experiments already referred to. I was slow in recognizing that it was of secondary importance, and not the main process of flexure. Judgment was confused by the existence of permanently warped specimens see § 4 below.

kilogram) it becomes ultimately comparable in amount with the elastic flexure

Temperature of baking, ° C	Permanent elongation, mm	Flexure by $\pm 1$ kilogram		Ratio inelastic total
		Total mm	Inelastic mm	
Not baked	0	2 3	0 16	0 07
55	0 015 ?	2 3	0 15	0 07
77	0 03	2 6	0 28	0 11
114	0 04	3 1	0 35	0 11
144	0 14	3 9	0 50	0 13
182	0 27	5 5	0 91	0 17
221	0 45	7 7	1 7	0 22
266	0 71	11 0	2 7	0 24
297	0 94	16 0	5 3	0 33
336	1 29	25 8	10 3	0 40
365	1 56	Broke		

While the table gives the flexure, elastic and permanent, due to the full load of  $\pm 1$  kilogram, it does not show the effect of a progressive increase or diminution of the load. This is conveniently exhibited by a cyclic hysteresis curve, similar in form to the curve of magnetic induction. A selection of three of these stress strain diagrams are shown in fig 3, which illustrate the progressive steepening and still more marked opening out of the diagram as the baking temperature is increased. To avoid confusion only the initial and final conditions, with one intermediate condition, are included.

It will be noticed that the marble in its initial condition, as received from the marble merchants, is capable of some inelastic flexure. The amount is small, only 0 16 mm under the standard load, and would not come into evidence without special arrangements, but there is no reason to doubt that it is genuine. Inadequate holding might simulate this effect, but, as already explained, this source of error was tested for and excluded.

Other untreated samples of marble show this effect, and sometimes much more conspicuously. A piece closely similar to the one used for the main series of tests gave an inelastic flexure of 0 38 mm under a load of  $\pm 500$  grams, it was thus considerably more flexible than the foregoing.

I had also a sample of pale grey (dove) marble from Messrs Wilkins, which they had noticed to be flexible. It was untreated by heat, and was as imported from Italy. Unfortunately it broke under the load I applied before any definite tests had been made. Evidently in this specimen mechanical weakness goes with flexibility, as in the heat-treated specimen.

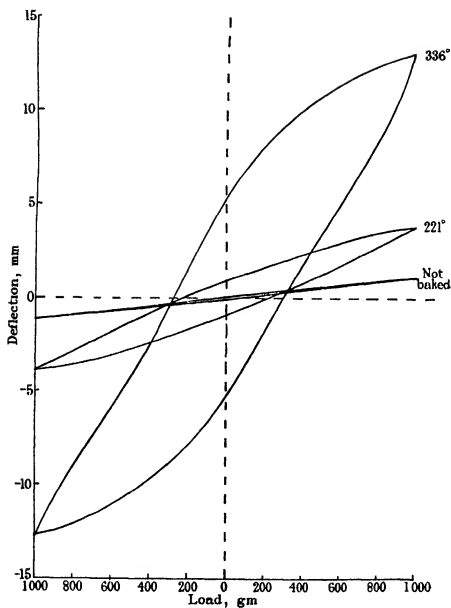


FIG 3.

These cases of natural flexibility in untreated marble deserve further investigation. In particular, it would be important to determine whether the specimen may not have got heated in the process of sawing it up.

### § 3 *General Discussion of Flexibility*

The marble used in these experiments consists of calcite crystal grains which, as viewed in a thin section under the microscope seem to fill practically the whole space. The size of these calcite grains is about 0.1 mm. and the edges are not infrequently fairly straight. It is difficult to make out any very definite evidence of cementing material between the grains, and it seems possible that they are simply adhering by close contact\*. It would seem at all events that the structure cannot hold together as it does without some adhesion between the grains, for the individual grain have no obvious re-entrant angles, and therefore cannot be held together simply by interlocking.

Now calcite is known to expand by heat much more in the direction of the optic axis than in the perpendicular direction. Indeed this is an understatement for in the latter direction the thermal expansion is actually negative†. The rhombohedron of calcite may be regarded as derived from a cube by shortening one of the diagonals, the edges which run up to one end of this diagonal becoming obtuse instead of right-angled. Thermal expansion tends to reverse this process and to diminish these obtuse angles. Mitscherlich found, in fact, that the rhombohedral angle was  $105^{\circ} 4'$  at  $10^{\circ} \text{C}$ , but was reduced to  $104^{\circ} 56'$  at  $110^{\circ} \text{C}$ ‡. The actual grains are not, of course, of rhombohedral shape, but their irregular shape similarly alters with temperature, and a grain which fitted accurately into the crystalline mosaic at ordinary temperature is no longer able to do so when the temperature is raised. The result of this is a springing apart of the contiguous grains, necessarily involving a partial rupture of the adhesion between them. In this way the whole structure is loosened, and progressive loss of mechanical rigidity results. The known permanent elongation of marble§ as recorded in the table, p. 272, is in accordance with this.

\* Mr F. Twyman, F.R.S., has kindly informed me in answer to a question that it is possible though not easy for the practical optician to put flat calcite surfaces into adhesive optical contact.

† Benoit's values for the coefficient of expansion are parallel to axis  $2.4963 \times 10^{-6}$  and perpendicular to axis  $-5.541 \times 10^{-6}$ . These are in agreement with the earlier work of Fizeau. See Tutton, "Crystallography and Crystal Measurement," vol. 2, p. 1329.

‡ See Tutton, vol. 2, p. 1303.

§ The permanent expansion of marble is described by Schäd, 'Phys. Rev.', vol. 10, p. 74 (1917).

general point of view and may be said decidedly to confirm it for the two effects become sensible at about the same temperature

Two other indications of the general loosening of the structure may be mentioned First, it is found that ink will "run" on marble that has been baked much more freely than on marble in its original condition Fig 4, Plate 13, shows the effect for indian ink, and fig 5, Plate 13, for ordinary writing ink, which flows more freely than indian ink In each of these pictures the marble on the left was baked for 24 hours at  $365^{\circ}\text{C}$  That on the right was a piece of the same, not baked The lines are ruled with an ordinary drawing pen, to ensure equal treatment a line was drawn on each piece of marble alternately without any refilling or readjustment of the pen throughout The running of the ink is, of course, attributed to the increased porosity of the material (*cf* blotting paper and writing paper)

Secondly, it is noticeable at the higher temperatures, marble is extremely friable, and can easily be crumbled with the help of pliers, or even broken off with the fingers No measurements have been made on the ultimate strength after baking at various temperatures Such measurements would probably not be very definite, but the observations casually made suggest a progressive loss of ultimate strength, proceeding *pari passu* with the elongation and increased flexibility It is this which sets a limit to the temperature of baking (see table, p 272)

At a high enough temperature marble, of course, loses carbon dioxide, and is converted into quicklime I was at one stage disposed to regard this change as conditioning the increased flexibility and tendency to crumble, but the experiments show marked increase of flexibility at temperatures below  $150^{\circ}\text{C}$ , which would seem to be too low a temperature for serious loss of carbon dioxide to take place A special experiment clears up any doubt A strip of marble was weighed as carefully as the rough balance available would allow, baked in the kitchen oven to a high degree of flexibility (temperature not determined), and weighed again The weight was 425 grams No loss of weight was indicated, and the loss, if any, was certainly less than 0.2 gram. It is clear, therefore, that there was no appreciable loss of carbon dioxide

Although the diminished rigidity of marble is generally accounted for on the above lines, more must be said to fill in the details We have to explain not only increased elastic flexure, but also the inelastic flexure which increases in relative importance the more the structure is loosened As to the increased elastic flexure the number of grain-contacts at which the internal stress is

taken must be diminished, and the stress at each increased, producing an increased strain

As regards inelastic flexure, it is natural to turn in the first instance to the example of other crystalline aggregates, and metals are the most obvious. In these, inelastic flexure occurs by the individual crystalline grains changing their shape by the occurrence of slip or twinning on particular crystallographic planes, and the work of Adams and Nicholson\* indicates that in the flow of marble under stresses of the order of 18,000 lbs per square inch the process is in many respects similar to that in metals. But in the present case the extreme forces to which the marble is subjected are only of the order of 450 lbs per square inch.†

Both the elastic and the inelastic flexure of baked marble are to most observers very surprising and unexpected effects. Considering the way that they develop together as the baking temperature is raised (table, p 272) it seems probable that they are closely connected. For reasons already given, the increased elastic flexure is connected with the loosening of contacts between adjacent grains, and if this is correct for the inelastic also, then any comparison with the metals is altogether wide of the mark. It seems that a plausible explanation can be derived if we combine the idea of elastic flexure with that of solid friction. The latter is supplied by the motion of the partially loosened grains one over the other. When the bending force is removed, this friction prevents complete recovery of straightness, by opposing the elastic force of restitution. The more the structure is loosened the greater becomes the friction and the less the elastic restoring force, and we are led to anticipate the inelastic bending becoming a progressively greater fraction of the total, as observed. If the structure is thought of as approaching complete disintegration, the friction would become all important, and the whole bending would approach to being inelastic. For obvious reasons we cannot approach very near the extreme case experimentally.

According to the point of view explained, the elastic restoring force, and the friction which prevents it from fully asserting itself, are considered to exist alongside of one another in the structure. As a rough model which may assist the imagination, consider a hinge working with some friction, and uniting two

\* *Loc cit*, p 372

† This figure is derived by treating the bent marble as perfectly elastic, so that the stress is proportional to the distance from the neutral axis. This supposition cannot, of course, be exact where Hook's law is not followed, but alternative suppositions, such as that the whole stress due to bending is uniformly taken on an outside skin of 1.5 mm thickness would lead to a similar result.



long arms. Let these be straightened out, and let a flexible steel lath be tied alongside. Then, on bending the model and releasing it again, the friction of the hinge will prevent the full recovery of straightness. The effect of baking the marble at a higher temperature will be represented by making the elastic lath thinner. This will diminish the force required to bend it, and increase the amount of permanent flexure.

It has been pointed out to me that the observed flexure of unbaked marble in the above experiments leads to a value of Young's modulus much less than that found for single calcite crystals. These give a value of the modulus between  $5.8 \times 10^{11}$  and  $11 \times 10^{11}$  according to the direction of the specimen in relation to the crystal axes,\* whereas the data on p. 272 lead to a value of  $2.6 \times 10^{11}$ , and another sample, which I have examined, gave a value of only  $0.79 \times 10^{11}$ . From this it seems clear that the crystals are not firmly united all over as in a metal, and that in fact there is some looseness of structure even in untreated marble. The same may indeed also be inferred from the varied behaviour of different samples as regards elastic flexure, and from the fact that they are all to some extent capable of permanent flexure.

#### § 4 *Warping of Marble*

We have seen in the introduction that some bent specimens of marble are incapable of being completely straightened by the external application of force.

Permanent bending or warping may be attributed to unsymmetrical heating. The known permanent expansion, if it occurred at one side only, would necessarily have this result. To illustrate the matter experimentally, the top surface of a piece of marble was made the bottom of an electric oven, and the under surface was kept cool by immersion in a shallow trough of running water. It was found that as expected, the hot side became permanently convex, and the cool one concave. The length of one piece used was 40.8 cm., thickness 1.9 cm. Heated to  $300^{\circ}\text{C}$  on one side, a marked curvature was developed. The sagitta was 1.25 mm. and the radius consequently 16.7 metres.

Such bent slabs are not flexible owing to the rigidity of the side that has not been heated. They imitate the bent piece of unknown origin which was mentioned in the introductory section, and maintain their curvature when placed with the convex side up and heavily loaded. Two such pieces, placed with the concave faces together, are shown in fig. 6, Plate 13.

\* Voigt, 'Ann. Physik,' vol. 39, p. 412, (1890). Quoted by Landolt and Bornstein, Tabellen, vol. 1, p. 84.

Unilateral heating has already been indicated by Kiestringer\* as a probable cause of the deformation of marble mantelpieces. As we have seen, however, these may become sufficiently baked right through to acquire flexibility. In the present experiment, the baking right through is prevented by water cooling, and the curvature is very great compared with that ordinarily seen in mantelpieces.

The curvature developed by unilateral heating is of the order of magnitude to be expected from the permanent expansion on the convex side. Let us take the piece above mentioned. With length  $l$ , expansion  $d l$ , and thickness  $t$ , we should have

$$\frac{l}{d l} = \frac{r}{t} = \frac{1870}{1.9} = 880$$

On the actual length of 40.8 cm. this would mean an expansion of 0.465 mm. Actually the expansion of the uniformly baked strip for 300° C. was somewhat larger than this, about 1 mm., see table, p. 272. Unfortunately the warped piece of marble did not come from the same source as the uniformly baked piece with which it is being compared. But even if they had been of identical texture, we must remember that with unilateral heating, while the temperature gradient may be regarded as uniform through the section, the permanent length increases much more rapidly at the higher temperatures (see p. 272). Thus the normal expansion ( $\epsilon$ ), that which would take place in the absence of constraint, is not able fully to assert itself on the convex side. The warped piece is presumably in a state of stress, the convex side being in compression, and the concave in tension.

### § 5 Possible Geological Application

The possible importance for geology of the permanent change of marble on heating should not be overlooked. By examining a laboratory specimen we can determine a limit to the temperature to which it can have been heated since the time when the calcite grains were formed into a compact aggregate as at present. Applying this criterion to the Carrara marble examined, the temperature cannot have risen so high as 77° C., for if this had occurred the specimen should be incapable of showing the elongation and loss of mechanical stiffness which it does show when baked at this temperature. But caution is perhaps necessary in applying such conclusions to material which has been at great pressures deep down in the earth's crust.

\* "Umschau," vol. 35, p. 436 (1931)



FIG. 5

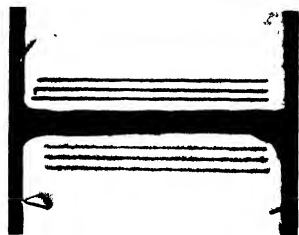


FIG. 4



FIG. 6



### § 6. *Summary*

Marble which has been baked at a temperature of  $100^{\circ}\text{C}$  or even less has its rigidity diminished. This effect becomes increasingly conspicuous up to temperatures of  $350^{\circ}\text{C}$ , when the bending under a given force may become 10 times as great as at first. Stress-strain diagrams are given showing hysteresis, the flexure being partly elastic, partly permanent. The progressive change of mechanical properties is traced as the baking temperature is increased. At the same time the marble becomes permanently elongated, its porosity is increased, as evidenced by the spreading of ink lines ruled on it, and its ultimate strength is diminished.

These results are interpreted as due to the unequal expansion of calcite by heat. The expansion is a maximum along the optic axis, and actually negative in the perpendicular direction. Thus the shape of each grain changes on heating, and they are no longer able to fit into the initial closely packed mosaic. The structure is dislocated and loosened, with increase of volume, and this process is irreversible. It is probable that with the moderate forces acting in these experiments, the individual grains are not deformed by slip or twinning, as when metals or marble are caused to flow, but that their elastic deformation, combine with friction in the loosened structure, will account for everything. Experiments are described on the warping of marble by unilateral heating, which are also brought under the same general point of view.

### DESCRIPTION OF PLATE.

FIG 4 —Right. Lines on marble in initial condition. Left. Lines on marble baked at  $365^{\circ}\text{C}$ . Indian ink used. Actual size.

FIG 5 —The same. Writing ink used.

FIG 6 —Two strips of marble warped by heating the (eventually) convex faces to  $300^{\circ}\text{C}$ . and keeping the others at  $15^{\circ}\text{C}$ ., water cooled. The concave faces are placed together. One-third actual size.

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# *The Energies of Alpha and Gamma Rays*

By H. A. WILSON, F.R.S., Rice Institute, Houston, Texas.

(Received December 14, 1933)

The energies of disintegration of the radioactive atoms, calculated from the alpha ray energies, may be arranged in pairs having sums which are multiples of 3.85\*. The following table shows this for the disintegration energies of radium C'. The values of the disintegration energies are those given by Rutherford, Lewis, and Bowden† and the unit is  $10^6$  electron volts

## Pairs of Disintegration Energies. Multiples of 3.85

78.29 + 91.12 = 169.41	$3.85 \times 44 = 169.40$
78.29 + 94.93 = 173.22	$3.85 \times 45 = 173.25$
78.29 + 102.69 = 180.98	$3.85 \times 47 = 180.95$
91.12 + 106.26 = 196.38	$3.85 \times 51 = 196.35$
92.41 + 0 = 92.41	$3.85 \times 24 = 92.40$
94.93 + 105.26 = 200.19	$3.85 \times 52 = 200.20$
96.73 + 99.68 = 196.41	$3.85 \times 51 = 196.35$
96.73 + 103.42 = 200.15	$3.85 \times 52 = 200.20$
100.97 + 107.09 = 208.06	$3.85 \times 54 = 207.90$
102.69 + 105.26 = 207.95	$3.85 \times 54 = 207.90$

The sum of all these pairs is 1825.16, which is equal to  $3.8506 \times 474$

The following table shows the same thing for pairs of disintegration energies of the other radioactive atoms. The values used are those given by Rutherford, Wynn-Williams, Lewis, and Bowden.†

## Pairs of Disintegration Energies Multiples of 3.85.

Rn, AcA	55.80 + 75.08 = 130.88	$3.85 \times 34 = 130.90$
Rn, AcCa	55.80 + 67.39 = 123.19	$3.85 \times 32 = 123.20$
RaCa, RaCa <sub>2</sub>	56.15 + 55.52 = 111.67	$3.85 \times 29 = 111.65$
RaCa <sub>2</sub> , AcC'	55.53 + 63.83 = 119.35	$3.85 \times 31 = 119.35$
Ana <sub>2</sub> , AcC'	66.85 + 75.81 = 142.66	$3.85 \times 37 = 142.45$
ThA, ThC'	68.97 + 96.61 = 165.58	$3.85 \times 43 = 165.55$
ThC', ThC''	96.61 + 107.41 = 204.02	$3.85 \times 53 = 204.05$
ThCa <sub>2</sub> , ThC'	62.04 + 107.41 = 169.45	$3.85 \times 44 = 169.40$
Po, Ana <sub>2</sub>	54.06 + 65.31 = 119.37	$3.85 \times 31 = 119.35$
ThCa	61.64 + 0 = 61.64	$3.85 \times 16 = 61.60$
Ana	69.37 + 0 = 69.37	$3.85 \times 18 = 69.30$

The sum of these pairs is 1416.98, which is equal to  $3.8505 \times 368$ . The value 3.8505 agrees almost exactly with the value 3.8506 got from the pairs of radium C' energies.

\* A preliminary account of some of the results given in this paper has been given in a letter to the Editor of the 'Physical Review' ('Phya. Rev.' Nov. 15, 1933).

† 'Proc. Roy. Soc.,' A, vol. 142, p. 347 (1933).

‡ 'Proc. Roy. Soc.,' A, vol. 129, p. 617 (1933).

The following table gives similar pairs of disintegration energies, each pair containing one radium C' energy and an energy for another atom —

Pairs of Disintegration Energies. Multiples of 3 85

RaCa	55 52 + 98 44 = 153 96	3 85 × 40 = 154 00
AcC'	75 81 + 78 29 = 154 10	3 85 × 40 = 154 00
AcC'	75 81 + 105 26 = 181 07	3 85 × 47 = 180 95
Ana <sub>1</sub>	66 65 + 91 12 = 157 77	3 85 × 41 = 157 85
Ana <sub>2</sub>	66 65 + 94 93 = 161 58	3 85 × 42 = 161 70
Ana <sub>3</sub>	66 65 + 102 69 = 169 34	3 85 × 44 = 169 40
RaA	61 09 + 96 73 = 157 82	3 85 × 41 = 157 85
ThCa <sub>1</sub>	62 04 + 99 68 = 161 72	3 85 × 42 = 161 70
ThC'	96 61 + 84 37 = 180 98	3 85 × 47 = 180 95
ThC'	96 61 + 99 68 = 196 29	3 85 × 51 = 196 35
ThC'	62 04 + 107 41 = 169 45	3 85 × 44 = 169 40
ThCa	61 64 + 92 41 = 154 05	3 85 × 40 = 154 00
Ana	69 37 + 92 41 = 161 78	3 85 × 42 = 161 70

The sum of all these pairs is 2159 91, which is equal to  $3\ 8501 \times 561$ . The sum of all the pairs is 5402 05, which is equal to  $3\ 8504 \times 1403$ . Thus it appears that the above three sets of pairs give values of this constant which differ from the mean value  $3\ 8504$  by  $+0\ 0002$ ,  $+0\ 0001$ , and  $-0\ 0003$  respectively, or by less than one part in twelve thousand.

The relative values of the disintegration energies are believed to be accurate to about one part in five thousand, but the absolute values may be in error by as much as one part in one thousand.

The constant  $3\ 8504 \times 10^5$  electron volts appears to have very exactly the same value for all the radioactive atoms so that it must represent some general atomic property.

The gamma ray energies of the different radioactive atoms may also be arranged in pairs having sums which are equal to multiples of a constant which is nearly equal to 3 85. This was shown for radium C' by the writer in a paper recently communicated to the Royal Society.

The relative values of the gamma ray energies for any one substance are known in some cases to about one in one thousand, but the absolute values may be in error by several parts in one thousand.

The following table gives the values of the constant, multiples of which are equal to the sums of pairs of the gamma ray energies, for several radioactive bodies —

Substance	Authors of energy determinations	Value of constant
RaC'	Ellis and Skinner, 1924	3 89
RaB	Ellis and Skinner, 1924	3 88
ThB, ThC	Ellis, 1932	3 85
RaC'	Ellis, 1933	3 85
RaB	Ellis, 1933	3 83

It seems probable that the differences between these values and 3 8504 are due to experimental errors in the absolute values. Since the alpha ray energies are believed to be correct to within one part in one thousand it follows that the earlier values of the energies of the RaC' and RaB rays are too large by about 1%.

An exact determination of the energies of any alpha or gamma ray would enable the energies of all the others to be corrected so that the errors in the absolute energies would be no greater than those in the relative energies.

A determination of the energies of several beta rays is now in progress in the writer's laboratory, which it is hoped will give the absolute values to within one part in ten thousand.

The fact that the alpha and gamma ray energies can be arranged in pairs with sums equal to  $3\ 85n$  shows that the energies are equal to  $3\ 85n \pm c_m$ , where  $c_m$  has a number of different values. It has been shown by the writer\* that the  $c$ 's for the alpha rays are approximately equal to the  $c$ 's for radium C'. It is also found that the  $c$ 's can be expressed as sums of multiples of the electronic energy levels. The electronic energy levels for the different radioactive atoms do not differ much and the alpha ray energies are not known with sufficient accuracy to enable the small differences between the  $c$ 's, for alpha rays, due to the differences between the electronic energy levels of the different atoms to be detected.

A simple system of energy levels which explains the emission of gamma rays with energies  $3\ 85n \pm c_m$  is shown in fig. 1. Only four values of  $c_m$  are shown for simplicity.

The fact that the  $c$ 's are independent of  $n$  suggests that the energy levels belong to two separate or only slightly coupled systems and since the  $c$ 's are related in a simple way to the electronic energy levels it is natural to suppose that the two systems are the nucleus and the electrons as previously suggested by the writer.

According to this, then, we have the remarkably simple result that the possible energies of any radioactive nucleus are equal to  $3\ 85n$ , where  $n = 1, 2, 3,$

If a nucleus disintegrates emitting an alpha ray the disintegration energy must be  $3\ 85(n - n')$  because the energy before disintegration is  $3\ 85n$  and  $3\ 85n'$  after. In the same way when the nucleus emits a gamma ray the energy of the gamma ray will be equal to  $3\ 85(n - n')$ . The observed energies

\* 'Phys. Rev.', vol. 44, p. 858 (1933)



include the energy changes of the electronic system  $\pm c_m$ . The energy transfer may take place, at the moment of emission, between the electrons and the nucleus or between the particle emitted and the electrons. It is impossible at present to distinguish between the two possibilities, in fact, it is doubtful if there is any real difference between them.

The simplest supposition we can make to explain the possible energies  $3.85n$  is that the nucleus acts like an elastic sphere which can vibrate with a frequency  $\nu$  given by  $h\nu = 3.85 \times 10^6$  electron volts. The possible energies are then  $h\nu(n + \frac{1}{2})$ , if we suppose the vibrations are analogous to those of a

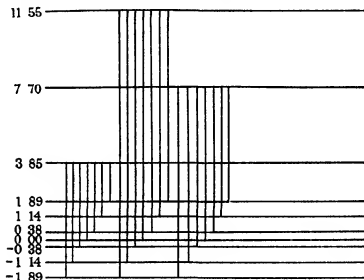


FIG. 1

one-dimensional harmonic oscillator. The frequency must be supposed independent of the mass and charge of the nucleus.

If a nucleus contains only protons and neutrons, then when it emits a beta ray a neutron must be converted into a proton. Possibly a gamma ray splits up into a beta ray and a positron which combines with a neutron. We should expect the beta ray to be emitted with energy  $3.85n$  like the alpha and gamma rays and the observed continuous energy distribution of the nuclear beta rays must be attributed to some secondary action such as collisions with the electrons\*. It may be significant that the average beta ray energy per dis-

\* The beta rays may either lose energy to or gain energy from the electrons.

integration of radium E is about  $3.9 \times 10^5$  electron volts, which is nearly equal to  $3.85 \times 10^5$

Another way of explaining the energies  $3.85n$  is to suppose that  $3.85$  is the energy set free when a neutron combines with a proton. The uranium nucleus may be supposed to contain 119 protons, 119 neutrons, and 27 electrons. The combination of the protons and neutrons would give energy  $119 \times 3.85 = 458$ . This is nearly equal to the total energy emitted in the uranium series of disintegrations which is about 45 million electron volts. The change of atomic weight corresponding to  $3.85 \times 10^5$  electron volts is 0.00041, and this is a possible value for the loss of atomic weight when a proton and a neutron combine to form a deuteron, but, of course, the masses of the protons and neutrons in a nucleus are not equal to their masses in the free state.

A theory of nuclear energy differences was suggested by Rutherford and Ellis\*. According to this theory there are two main energy levels which, to explain the energy differences  $3.85n \pm c_m$ , may be supposed here to differ by  $3.85$ . If there are  $p_1$  alpha particles in the upper level and  $p_2$  in the lower level, the energy is  $3.85p_1 + f(p_1, p_2)$ , where  $f(p_1, p_2)$  is a function of  $p_1$  and  $p_2$  which represents the energy of mutual action between the alpha particles. If  $n$  alpha particles drop from the upper to the lower level, the energy change

$$3.85n + f(p_1, p_2) - f(p_1 - n, p_2 + n)$$

The differences  $f(p_1, p_2) - f(p_1 - n, p_2 + n)$  would then represent the  $c$ 's but we should expect these differences to depend on  $n$ , whereas the  $c$ 's in  $3.85n \pm c_m$  are independent of  $n$ . This theory, therefore, does not seem to be capable of explaining energy differences  $3.85n \pm c_m$  with  $c$ 's independent of  $n$ .

In a recent paper by Rutherford, Lewis, and Bowden† a scheme of energy levels for the radium C' nucleus is proposed. This scheme gives a large number of the gamma rays, but it does not seem as well fitted to represent energy differences equal to  $3.85n \pm c_m$  as the simple scheme suggested above.

The nuclear energy constant  $3.85 \times 10^5$  electron volts is equal to  $6.122 \times 10^7$  ergs. The corresponding wave-length given by  $6.122 \times 10^{-7} = hc/\lambda$  is equal to  $3.21 \times 10^{-10}$  cm. The corresponding wave-number is  $3.12 \times 10^9$

\* 'Proc. Roy. Soc.,' A, vol. 132, p. 667 (1931)

† 'Proc. Roy. Soc.,' A, vol. 142, p. 347 (1933)

It seems clear that this constant represents some fundamental atomic quantity, the nature of which is as yet uncertain

*Summary*

It is shown that the energies of disintegration of the radioactive atoms may be arranged in pairs having sums equal to multiples of  $3.8504 \times 10^5$  electron volts. This constant must, therefore, represent some general atomic property, the nature of which is as yet uncertain

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*Two New Properties of Mathematical Likelihood.*

By R. A. FISHER, F.R.S.

(Received December 14, 1933)

1 *Introductory*

To Thomas Bayes\* must be given the credit of broaching the problem of using the concepts of mathematical probability in discussing problems of inductive inference, in which we argue from the particular to the general, or, in statistical phrasology, argue from the sample to the population, from which, *ex hypothesi*, the sample was drawn. Bayes put forward, with considerable caution, a method by which such problems could be reduced to the form of problems of probability. His method of doing this depended essentially on postulating *a priori* knowledge, not of the particular population of which our observations form a sample, but of an imaginary population of populations from which this population was regarded as having been drawn at random. Clearly, if we have possession of such *a priori* knowledge, our problem is not properly an inductive one at all, for the population under discussion is then regarded merely as a particular case of a general type, of which we already possess exact knowledge, and are therefore in a position to draw exact deductive inferences.

To the merit of broaching a fundamentally important problem, Bayes added that of perceiving, much more clearly than some of his followers have done, the logical weakness of the form of solution he put forward. Indeed we

\* 'Phil. Trans.,' vol. 53, p. 370 (1763)

are told that it was his doubts respecting the validity of the postulate needed for establishing the method of inverse probability that led to his withholding his entire treatise from publication. Actually it was not published until after his death.

If a sample of  $n$  independent observations each of which may be classified unambiguously in two alternative classes as "successes" and "failures" be drawn from a population containing a relative frequency  $x$  of successes, then the probability that there shall be  $a$  successes in our samples is, as was first shown by Bernoulli,

$$\frac{n!}{a!(n-a)!} x^a (1-x)^{n-a} \quad (1)$$

This is an inference, drawn from the general to the particular, and expressible in terms of probability. We are given the parameter  $x$ , which characterizes the population of events of which our observations form a sample, and from it can infer the probability of occurrence of samples of any particular kind.

If, however, we had *a priori* knowledge of the probability,  $f(x) dx$ , that  $x$  should lie in any specified range  $dx$ , or if, in other words, we knew that our population had been chosen at random from the population of populations having various values of  $x$ , but in which the distribution of the variate  $x$  is specified by the frequency element  $f(x) dx$  of known form, then we might argue that the probability of first drawing a population in the range  $dx$ , and then drawing from it a sample of  $n$  having  $a$  successes, must be

$$\frac{n!}{a!(n-a)!} x^a (1-x)^{n-a} f(x) dx, \quad (2)$$

since this sequence of events has occurred for some value of  $x$ , the expression (2) must be proportional to the probability, subsequent to the observation of the sample, that  $x$  lies in the range  $dx$ . The postulate which Bayes considered was that  $f(x)$ , the frequency density in the hypothetical population of populations, could be assumed *a priori* to be equal to unity.

As an axiom this supposition of Bayes fails, since the truth of an axiom should be manifest to all who clearly apprehend its meaning, and to many writers, including, it would seem, Bayes himself, the truth of the supposed axiom has not been apparent. It has, however, been frequently pointed out that, even if our assumed form for  $f(x) dx$  be somewhat inaccurate, our conclusions, if based on a considerable sample of observations, will not greatly be affected, and, indeed, subject to certain restrictions as to the true form of  $f(x) dx$ , it may be shown that our errors from this cause will tend to zero as the

sample of observations is increased indefinitely. The conclusions drawn will depend more and more entirely on the facts observed, and less and less upon the supposed knowledge *a priori* introduced into the argument. This property of increasingly large samples has been sometimes put forward as a reason for accepting the postulate of knowledge *a priori*. It appears, however, more natural to infer from it that it should be possible to draw valid conclusions from the data alone, and without *a priori* assumptions. If the justification for any particular form of  $f(x)$  is merely that it makes no difference whether the form is right or wrong, we may well ask what the expression is doing in our reasoning at all, and whether, if it were altogether omitted, we could not without its aid draw whatever inferences may, with validity, be inferred from the data. In particular we may question whether the whole difficulty has not arisen in an attempt to express in terms of the single concept of mathematical probability, a form of reasoning which requires for its exact statement different though equally well-defined concepts.

If, then, we disclaim knowledge *a priori*, or prefer to avoid introducing such knowledge as we possess into the basis of an exact mathematical argument, we are left only with the expression

$$\frac{n!}{a!(n-a)!} x^a (1-x)^{n-a},$$

which, when properly interpreted, must contain the whole of the information respecting  $x$  which our sample of observations has to give. This is a known function of  $x$ , for which, in 1922, I proposed the term "likelihood," in view of the fact that, with respect to  $x$ , it is not a probability, and does not obey the laws of probability, while at the same time it bears to the problem of rational choice among the possible values of  $x$  a relation similar to that which probability bears to the problem of predicting events in games of chance. From the point of view adopted in the theory of estimation, it could be shown, in fact, that the value of  $x$ , or of any other parameter, having the greatest likelihood possessed certain unique properties, in which such an estimate is unequivocally superior to all other possible estimates. Whereas, however, in relation to psychological judgment, likelihood has some resemblance to probability, the two concepts are wholly distinct, in that probability is appropriate to a class of cases in which uncertain inferences are possible from the general to the particular, while likelihood is appropriate to the class of cases arising in the problem of estimation, where we can draw inferences, subject to a different kind of uncertainty, from the particular to the general.

The primary properties of likelihood in relation to the theory of estimation have been previously demonstrated \*. In the following sections I propose to exhibit certain further properties arising when the functional properties of the specification of the population fulfil certain special, but practically important, conditions

## 2 *The Distribution of Sufficient Statistics*

The essential feature of statistical estimates which satisfy the criterion of sufficiency is that they by themselves convey the whole of the information, which the sample of observations contains, respecting the value of the parameters of which they are sufficient estimates. This property is manifestly true of a statistic  $T_1$ , if for any other estimate  $T_2$  of the same parameter,  $\theta$ , the simultaneous sampling distribution of  $T_1$  and  $T_2$  for given  $\theta$ , is such that given  $T_1$ , the distribution of  $T_2$  does not involve  $\theta$ , for if this is so it is obvious that once  $T_1$  is known, a knowledge of  $T_2$ , in addition, is wholly irrelevant, and if the property holds for all alternative estimates, the estimate  $T_1$  will contain the whole of the information which the sample supplies

This remarkable property will be possessed when

$$\frac{\partial}{\partial \theta} \log L,$$

where  $L$  is the likelihood of  $\theta$  for a given sample of observations, is the same function for all samples yielding the same estimate  $T_1$ , for on integrating the expression above with respect to  $\theta$ , it appears that  $\log L$  is the sum of two components, one a function only of  $\theta$  and  $T_1$ , and the other dependent on the sample but independent of  $\theta$ . If

$$f(T_1, T_2, \theta) dT_1 dT_2$$

is the frequency with which samples yield estimates simultaneously in the ranges  $dT_1$  and  $dT_2$ , it follows that

$$f(T_1, T_2, \theta) = \phi_1(T_1, \theta) \phi_2(T_1, T_2),$$

where the first factor involves  $T_1$  and  $\theta$  only, and the second does not involve  $\theta$ . The distribution of  $T_2$  given  $T_1$  will therefore be

$$\phi_2(T_1, T_2) dT_2 / \int \phi_2(T_1, T_2) dT_2$$

\* Fisher, 'Proc Camb Phil Soc,' vol 22, p 700 (1925)

the integral being taken over all possible values of  $T_s$ , and in this expression the parameter  $\theta$  is seen not to appear

The condition that  $\partial L/\partial \theta$  should be constant over the same sets of samples for all values of  $\theta$ , which has been shown to establish the existence of a sufficient estimate of  $\theta$ , thus requires that the likelihood is a function of  $\theta$ , which, apart from a factor dependent on the sample, is of the same form for all samples yielding the same estimate  $T$ . The sufficiency of sufficient statistics may thus be traced to the fact that in such cases the value of  $T$  itself alone determines the form of the likelihood as a function of  $\theta$ . If a conventional value such as unity is given to the maximum likelihood for any sample, the likelihood is thus expressible as a function of  $\theta$  and  $T$  only, if  $T$  is the sufficient estimate. We shall use this property in obtaining a general form for the sampling distribution of sufficient statistics.

2.1 It will help if we take an illustrative example of this problem. Let the element of frequency in a distribution be given by

$$\frac{1}{\theta!} x^\theta e^{-x} dx$$

where the variate  $x$  can take any real value from 0 to  $\infty$ , and  $\theta$  is an unknown parameter greater than  $-1$ . Consider the problem of estimating  $\theta$  from a sample of  $n$  values of  $x$ .

If  $L$  is the likelihood of any possible value of  $\theta$ ,

$$\log L = -n \log \theta! + \theta S(\log x) - S(x),$$

and this is maximized for variations of  $\theta$ , when  $\theta = T$ , where

$$F(T) = \frac{1}{n} S(\log x),$$

and  $F(T)$  is the first differential of the logarithm of the factorial function. This is the equation of estimation by the method of maximum likelihood. It will be observed that apart from a constant factor the likelihood is expressible as a function of  $\theta$  and  $T$  only, that is

$$L = A \exp \{-n \log \theta! + n \theta F(T)\}$$

so that  $T$  is evidently a sufficient estimate.

2.2 The sampling distribution of our estimate must evidently be derived from that of the mean of the logarithms of the several values of  $x$  in the sample. Now the mean value of

$$\frac{x}{\theta!} \log x$$

is

$$\int \frac{1}{\theta!} x^{\theta + \frac{u}{n}} e^{-x} dx = \frac{(\theta + \frac{u}{n})!}{\theta!}$$

By the familiar process of expanding the multiple integral in a product of single integrals, the mean value over all samples of

$$e^{\frac{u}{n} \sum \log x} = e^{u F(T)}$$

is

$$M = \frac{\left\{ \left( \theta + \frac{u}{n} \right)! \right\}^n}{\{\theta!\}^n}$$

and this is the characteristic function of

$$F(T)$$

from which its distribution may be inferred

To determine the probability function knowing the characteristic function  $M(u)$  we may use the property of the sine integral

$$\int_0^\infty \frac{\sin u}{u} du = \frac{\pi}{2},$$

writing  $ku$  for  $u$  it appears that

$$\frac{1}{\pi} \int_0^\infty \sin kt \frac{dt}{t} = \frac{1}{2}$$

when  $k$  is positive, and  $-\frac{1}{2}$  when  $k$  is negative, or that

$$\frac{1}{\pi} \int_0^\infty \{\sin(x-a)t - \sin(x-b)t\} \frac{dt}{t}$$

where  $b > a$ , is unity when  $b > x > a$ , and zero when  $x$  is less than  $a$  or exceeds  $b$ .

Consequently the Stieltjes integral

$$\int_a^b df(x) = \frac{1}{\pi} \int_0^\infty \frac{dt}{t} \int_{-\infty}^\infty \{\sin(x-a)t - \sin(x-b)t\} df(x),$$

writing

$$\sin(x-a)t = \frac{1}{2i} (e^{it(x-a)} - e^{-it(x-a)})$$

gives us

$$\int_a^b df(x) = \frac{1}{2\pi} \int_0^\infty \frac{dt}{t} \{e^{-iat} M(u) - e^{iat} M(-u) - e^{-ibt} M(u) + e^{ibt} M(-u)\},$$



where  $f(x)$  is the probability function of the variate  $x$ , and  $M$  is its characteristic function

We may note that  $M(u)$  and  $M(-u)$  must be conjugate quantities, which may be written  $R \pm iI$ , then

$$-e^{-iut} M(u) + e^{iut} M(-u) = 2iR \sin bt - 2iI \cos bt$$

so that the integral takes the real form

$$\frac{1}{\pi} \int_0^\infty \frac{dt}{t} \{R (\sin bt - \sin at) - I (\cos bt - \cos at)\}$$

Where the probability function is differentiable so that

$$df(x) = y dx$$

then

$$\begin{aligned} df(x) = y dx &= \frac{dx}{2\pi} \int_0^\infty \{e^{-iut} M(u) + e^{iut} M(-u)\} dt \\ &= \frac{dx}{\pi} \int_0^\infty \{R \cos (tx) + I \sin (tx)\} dt \end{aligned}$$

24 For the sufficient statistic  $T$ , the sampling distribution will therefore be given by

$$df = \frac{d^F(T)}{2\pi} \int_{-\infty}^\infty e^{-iuf(T)} M(u) dt,$$

but

$$e^{-iuf(T)} M' = \frac{L(\theta)}{L\left(\theta + \frac{u}{n}\right)},$$

hence the distribution may be directly inferred from the nature of the likelihood function in the form

$$df = \frac{F(T)}{2\pi} \frac{dT}{2\pi} \int_{-\infty}^\infty \frac{L(\theta)}{L\left(\theta + \frac{u}{n}\right)} dt$$

or

$$\frac{n}{2\pi} F(T) dT \int_{-\infty}^\infty \frac{L(\theta)}{L\left(\theta + \frac{u}{n}\right)} dt$$

where  $F(T)$  stands for the second differential coefficient of  $\log(T')$

We may illustrate the use of this formula by deriving the limiting forms for extreme values of  $\theta$

Near the limit  $\theta \rightarrow -1$  the general expression

$$\frac{n d(F(T))}{2\pi} \int_{-\infty}^\infty \frac{(\theta + u)^n}{\theta^n} e^{-iuf(T)} dt$$

may be rewritten, substituting

$$T = -1 + e^{-\theta}$$

$$\theta = -1 + e^{-\gamma},$$

and, since, as will be apparent, when  $\gamma$  is large, all important contributions will arise from small values of  $t$ , the limiting form of our distribution is

$$\frac{ne^g}{2\pi} \int_{-\infty}^{\infty} \left(1 + \frac{it}{1 + \theta}\right)^{-n} e^{itns^g} dt$$

Writing  $t$  for  $te^{\gamma}$  we then have

$$\frac{n}{2\pi} \frac{dq}{e^{(g-\gamma)}} \int_{-\infty}^{\infty} (1 + it)^{-n} e^{itns^{(g-\gamma)}} dt$$

and writing  $z$  for  $1 + it$

$$e^{-n\gamma} \frac{n}{2\pi i} \int z^{-n} e^{nuz} dz$$

where  $u$  stands for  $e^{g-\gamma}$ , and the integral is taken from  $1 - i\infty$  to  $1 + i\infty$  or in an open contour passing counter clockwise to the right of the singularity,  $z = 0$ . Writing  $\zeta$  for  $nuz$  we have

$$\frac{n(nu)^{n-1} e^{-n\gamma}}{2\pi i} \int \zeta^{-n} e^{\zeta} d\zeta,$$

where the integral does not now involve the variate  $u$ , and is evidently  $2\pi i/(n-1)!$ . The distribution now appears as

$$\frac{1}{(n-1)!} n^n u^{n-1} e^{-nu} du,$$

in which  $e^{g-\gamma}$  may be substituted for  $u$ .

The probability integral in this case is given by the  $\chi^2$  distribution for  $2n$  degrees of freedom. Thus if a sample of 10 had been taken, we have 20 degrees of freedom, and the 5% values are at  $\chi^2 = 10.851$  and  $31.410$ \*. Putting  $nu = \frac{1}{2}\chi^2$  the 5% values of  $u$  are 0.5426 and 1.5705, whence those of  $g - \gamma$  can be obtained, showing that in 90% of samples  $g$  will lie between  $\gamma - 0.6110$  and  $\gamma + 0.4514$ . For given  $g$ , therefore, the fiducial probability is 5% that  $\gamma$  exceeds  $g + 0.6110$ , or falls short of  $g - 0.4514$ .

At the upper limit where  $\theta \rightarrow \infty$  we may write

$$T = g^2, \quad \theta = \gamma^2,$$

\* "Statistical Methods for Research Workers," Table III

and since  $g - \gamma$  remains finite for finite values of the probability function

$$\log \frac{T}{\theta} = \frac{2(g - \gamma)}{\gamma}$$

and tends to zero The general expression for the distribution, which is

$$\frac{n d(F(T))}{2\pi} \int_{-\infty}^{\infty} \frac{(\theta + it)^n}{\theta^n} e^{-itnF(T)} dt,$$

tends to the limiting form

$$df = \frac{n dT}{2\pi T} \int_{-\infty}^{\infty} (t^{in} e^{-\frac{n^2}{2\sigma} T - itn} dt$$

or substituting for  $T$  and  $\theta$

$$\begin{aligned} df &= \frac{n dg}{\pi g} \int_{-\infty}^{\infty} e^{-2itn(g-\gamma)/\gamma} e^{-n^2/2\gamma^2} dt \\ &= \sqrt{\frac{2n}{\pi}} e^{-2n(g-\gamma)^2/\gamma^2} dg \end{aligned}$$

showing that  $g$  tends in the limit to be normally distributed about the population value  $\gamma$  with variance  $1/4n$  The 5% points of the distribution of  $g$  are therefore  $\gamma \pm 1.645/\sqrt{4n}$ , and for a given  $g$  the 5% points of the fiducial distribution of  $\gamma$  are  $g \pm 1.645/\sqrt{4n}$ \*

2.5 The interest of this form lies in the possibility of generalizing it for all sufficient statistics For, let the equation of maximum likelihood have a solution

$$\phi(T) = A$$

where  $A$  is a symmetric function of the observations not involving the parameter  $\theta$  The expression for  $\partial/\partial\theta \log L$  must have been of the form

$$C\{A\psi'(\theta) - \phi(\theta)\psi'(\theta)\}$$

where the possible factor  $C$ , if not a constant, must be a function of the observations which is expressible as a function of  $A$ , if the likelihood is to be expressible as a function of  $\theta$  and  $T$  only

The expression for  $\log L$  then must be of the form

$$CA\psi(\theta) - C \int \phi(\theta) d\psi(\theta) + B$$

\* "Statistical Methods for Research Workers," Table I

where  $B$  is a function of the observations only. That  $C$  a symmetric function of all the observations must be merely the number  $n$  in the sample appears from the fact that  $\log L$  is the sum of expressions involving each observation singly. Hence

$$CA = S(X), \quad B = S(X_1),$$

where  $X, X_1$ , are functions of the individual observations  $x$ . The likelihood is now the product

$$e^{-n \int \phi(\theta) d\phi(\theta)} e^{\phi(\theta) S(X)} e^{N(\lambda_1)}$$

and

$$\frac{L(\psi)}{L(\psi + u)} = e^{-u S(X)} e^{n F_1(\psi + u) - n F_1(\psi)}$$

where  $F_1(\psi)$  is written for  $\int \phi d\psi$

But the frequency function of the variate  $X$  was given by

$$e^{-F_1(\psi)} e^{\lambda \psi} e^{\lambda_1} \frac{dx}{dX},$$

hence its characteristic function is

$$M(u) = e^{F_1(\psi + u) - F_1(\psi)}$$

while that of  $S(X)$  is the  $n$ th power of this expression, hence the probability that  $S(X)$  lies between  $S_0$  and  $S_1$  is

$$\begin{aligned} \int_0^1 df &= \frac{1}{2\pi} \int_{-\infty}^{\infty} \frac{dt}{t} \{ e^{-i n t} M^n(u) - e^{i n t} M^n(-u) - e^{-i n t} M^n(u) + e^{i n t} M^n(-u) \} \\ &= \frac{1}{2\pi} \int_{-\infty}^{\infty} \frac{dt}{t} \left\{ \frac{L(\psi, S_0)}{L(\psi + u, S_0)} - \frac{L(\psi, S_0)}{L(\psi - u, S_0)} - \frac{L(\psi, S_1)}{L(\psi + u, S_1)} + \frac{L(\psi, S_1)}{L(\psi - u, S_1)} \right\} \end{aligned}$$

this being the general expression for the probability of any sufficient statistic falling within assigned limits,  $S_1$  and  $S_0$  being the limits of the known function  $n\phi(T)$  of the sufficient estimate  $T$ .

2.6 The property that where a sufficient statistic exists, the likelihood, apart from a factor independent of the parameter to be estimated, is a function only of the parameter and the sufficient statistic, explains the principal result obtained by Neyman and Pearson in discussing the efficacy of tests of significance. Neyman and Pearson introduce the notion that any chosen test of a hypothesis  $H_0$  is more powerful than any other equivalent test, with regard to an alternative hypothesis  $H_1$ , when it rejects  $H_0$  in a set of samples having

an assigned aggregate frequency  $\epsilon$  when  $H_0$  is true, and the greatest possible aggregate frequency when  $H_1$  is true

If any group of samples can be found within the region of rejection whose probability of occurrence on the hypothesis  $H_1$  is less than that of any other group of samples outside the region, but is not less on the hypothesis  $H_0$ , then the test can evidently be made more powerful by substituting the one group for the other. Consequently, for the most powerful test possible the ratio of the probabilities of occurrence on the hypothesis  $H_0$  to that on the hypothesis  $H_1$  is less in all samples in the region of rejection than in any sample outside it. For samples involving continuous variation the region of rejection will be bounded by contours for which this ratio is constant. The regions of rejection will then be required in which the likelihood of  $H_0$  bears to the likelihood of  $H_1$ , a ratio less than some fixed value defining the contour.

The test of significance is termed uniformly most powerful with regard to a class of alternative hypotheses if this property holds with respect to all of them. This evidently requires that the contours defined by the ratio of the likelihood of  $H_1$  and  $H_0$  shall be the same as those defined by the ratios of the likelihood of any two hypotheses in the class. If, therefore,  $T'$  is a statistic defining these contours, and  $\theta_1, \theta_2, \dots$  are variable parameters defining the hypothetical populations, the likelihood of any hypothesis must be expressed in the form

$$L = Af(T', \theta_1, \theta_2, \dots)$$

where  $A$  is a factor independent of the parameters

The method of estimation by maximum likelihood, when applied to the form above, will yield equations for  $\theta_1, \theta_2, \dots$ , etc

$$\begin{aligned}\phi_1(T', \theta_1, \theta_2, \dots) &= 0, \\ \phi_2(T', \theta_1, \theta_2, \dots) &= 0,\end{aligned}$$

where

$$\phi_i = \frac{\partial}{\partial \theta_i} \log f,$$

and the solutions of these will give estimates of  $\theta_1, \theta_2, \dots$ , which we may designate  $T_1, T_2, \dots$ , in the form

$$\begin{aligned}T_1 &= \psi_1(T') \\ T_2 &= \psi_2(T'), \text{ etc}\end{aligned}$$

It is evident, at once, that such a system is only possible when the class of hypotheses considered involves only a single parameter  $\theta$ , or, what comes to

the same thing, when all the parameters entering into the specification of the population are definite functions of one of their number. In this case, the regions defined by the uniformly most powerful test of significance are those defined by the estimate of maximum likelihood,  $T$ . For the test to be uniformly most powerful, moreover, these regions must be independent of  $\theta$ , showing that the statistic must be of the special type distinguished as sufficient. Such sufficient statistics have been shown to contain all the information which the sample provides relevant to the value of the appropriate parameter  $\theta$ . It is inevitable therefore that if such a statistic exists it should uniquely define the contours best suited to discriminate among hypotheses differing only in respect of this parameter, and it is surprising that Neyman and Pearson should lay it down as a preliminary consideration that "the testing of statistical hypotheses cannot be treated as a problem in estimation". When tests are considered only in relation to sets of hypotheses specified by one or more variable parameters, the efficacy of the tests can be treated directly as the problem of estimation of these parameters. Regard for what has been established in that theory, apart from the light it throws on the results already obtained by their own interesting line of approach, should also aid in treating the difficulties inherent in cases in which no sufficient statistic exists.

### 3 *A Second Class of Parameters for which Estimation need Involve no Loss of Information*

In the case of sufficient statistics the likelihood function is, apart from a constant factor, the same for all sets of observations which yield the same estimate by the method of maximum likelihood. A second case, of somewhat wider practical application, occurs when, although the sets of observations which provide the same estimate differ in their likelihood functions, and therefore in the nature and quantity of the information they supply, yet when samples alike in the information they convey exist for all values of the estimate and occur with the same frequency for corresponding values of the parameter.

The nature of the correspondence may be stated as follows. If  $x_1, \dots, x_n$  stands for a sample of  $n$  values of a variate  $x$ , the distribution of which is conditioned by a parameter,  $\theta$ , then for any value of  $\theta$ , there will be a definite probability

$$p(x, \theta)$$

of the occurrence of a variate less in value than  $x$ .

If, therefore, we take any other value of the parameter, say  $\phi$ , there will, with continuous variates, always exist a series of observational values,  $y$ , corresponding to the original series  $x$ , such that

$$p(y, \phi) = p(x, \theta)$$

The samples  $x$  and  $y$  will, however, only correspond in the sense required for our present purpose if corresponding to any possible value,  $\theta$ , a value,  $\phi$ , can be found so that the relationship above holds for all values of  $x$ . If, in fact, the equation were solved for  $y$ , in the form

$$y = f(x, \theta, \phi)$$

it is required that  $f$  shall be of the form

$$f(x, \theta, \phi) = \Omega(x, \theta) \quad (3)$$

where  $\Omega$  is a function of  $\theta$  and  $\phi$ , independent of the observations, and such that for any possible values of  $\theta$  and  $\Omega$  there exists a corresponding value of  $\phi$ . Stated symmetrically it is required that some function of  $x$  and  $y$  can be equated to a function of  $\theta$  and  $\phi$ .

The typical case of such a relationship occurs in parameters of location. If the distribution of the variate  $x$  involves a parameter  $\theta$ , such that the frequency with which  $x$  falls in any element  $dx$  of its range is a function of  $(x - \theta)$ , then  $\theta$  may be called a parameter of location. In such a case the functional relationship (3) may be written

$$x - y = \theta - \phi$$

and is clearly of the form required.

Let us take an example in which there is no sufficient estimate, and in which the loss of information in estimating the unknown parameter even by the method of maximum likelihood is considerable. The distribution of  $x$  is a double exponential curve, the probability of  $x$  falling in the range  $dx$ , being

$$\frac{1}{2} e^{-|x-\theta|} dx$$

The logarithm of the likelihood is

$$-S|x - \theta|,$$

and this increases when  $\theta$  is increased only if more observations are greater than those less than  $\theta$ . The likelihood is therefore maximized if the number of observations is odd, by equating  $\theta$  to the median observation, if the number

of observations is even the likelihood is constant when  $\theta$  has any value between the two central observations

For a sample of an odd number,  $n = 2s + 1$  of observations, the sampling distribution of the median is determinate, and the loss of information, if we use the median as an estimate, unsupplemented by the ancillary information which the sample contains, may be calculated. For, if the central observation lies at a distance  $u$  from the centre of the distribution,  $u$  being supposed positive, then the  $s$  highest values observed must each have fallen in a region comprising only

$$\frac{1}{2}e^{-u}$$

of the total frequency, while the  $s$  lowest values have fallen in the remaining region comprising

$$1 - \frac{1}{2}e^{-u}$$

of the total. Finally, the probability of the median itself falling in the range  $du$  is

$$\frac{1}{2}e^{-u} du,$$

so that compounding the independent probabilities into which the event has been analysed we have

$$df = \frac{(2s+1)!}{(s!)^2} \cdot (\frac{1}{2}e^{-u})^s (1 - \frac{1}{2}e^{-u})^s \frac{1}{2}e^{-u} du$$

as the probability of the median having a positive sampling error,  $u$ . As  $s$  is increased without limit we may write

$$u\sqrt{n} = t,$$

and the distribution tends to the limit

$$df = \frac{1}{\sqrt{2\pi}} e^{-\frac{1}{2}t^2} dt$$

The amount of information derivable from a large sample of  $n$  thus tends to equality with  $n$ , as the size of the sample is increased. Since the information supplied by the independent observations is additive,\* each must supply one unit, and a sample of  $2s + 1$  observations must contain  $2s + 1$  units of information. The quantity elicited by using the median,  $\frac{1}{2}e$ , by replacing the  $2s + 1$  observations from the distribution

$$df = \frac{1}{2} e^{-|x-\theta|} dx,$$

\* Fisher, 'Proc Camb Phil Soc' vol 22, p 700 (1925)



by a single observation from the distribution

$$df = \frac{(2s+1)!}{(s!)^2 2^{2s+1}} e^{-s|u-s|} (2 - e^{-|u-s|})^s e^{-|u-s|} du,$$

may be calculated from the mean value of

$$\left( \frac{d}{d\theta} \log \frac{df}{du} \right)^2,$$

or of

$$\left( s - \frac{se^{-|u-s|}}{2 - e^{-|u-s|}} + 1 \right)^2.$$

When  $s$  exceeds unity the average values may be evaluated from the consideration that

$$\int_0^\infty \frac{(2s+1)!}{(s-1)! (s+1)! 2^{2s+1}} e^{-|s-1|u} (2 - e^{-u})^{s-1} e^{-u} du$$

represents the probability that at least  $s+2$  observations have positive, and only  $s-1$  observations negative, deviations, and may therefore be equated to

$$\frac{1}{2} - \frac{(2s+1)!}{s! (s+1)! 2^{2s+1}}$$

The mean value of  $e^{-|u-s|}/(2 - e^{-|u-s|})$  is therefore found to be

$$\frac{s+1}{s} \left( 1 - \frac{(2s+1)!}{s! (s+1)! 2^{2s}} \right)$$

Similarly, the mean value of  $e^{2|u-s|}/(2 - e^{-|u-s|})^2$  is

$$\frac{(s+1)(s+2)}{s(s-1)} \left( 1 - \frac{(2s+1)!}{s! (s+1)! 2^{2s}} - \frac{(2s+1)!}{(s-1)! (s+2)! 2^{2s}} \right)$$

The amount of information provided by the median of  $2s+1$  observations is therefore

$$\begin{aligned} (s+1)^2 - 2s(s+1) &= \frac{s+1}{s} + s^2 \frac{(s+1)(s+2)}{s(s-1)} \\ &+ 2(s+1)^2 \frac{(2s+1)!}{s! (s+1)! 2^{2s}} - \frac{s(s+1)(s+2)}{s-1} \frac{(2s+1)!}{s! (s+1)! 2^{2s}} \\ &- \frac{s(s+1)(s+2)}{s-1} \frac{(2s+1)!}{(s-1)! (s+2)! 2^{2s}} \end{aligned}$$

or

$$\frac{(s+1)(2s+1)}{(s-1)} \left\{ 1 - \frac{(2s)!}{(s!)^2 2^{2s-1}} \right\}$$

In the special case,  $s = 1$ , the general method fails, and a direct integration yields the value

$$12 (\log 2 - \frac{1}{4}),$$

which is the limit to which the general expression tends as  $s \rightarrow 1$

The median is an efficient estimate in the sense of the theory of large samples, for the ratio of the amount of information supplied to the total available tends to unity as the sample is increased. Nevertheless, the absolute amount lost increases without limit. As  $s$  increases, this amount lost,

$$2s + 1 - \frac{(s+1)(2s+1)}{s-1} \left\{ 1 - \frac{2s!}{(s!)^2 2^{2s-1}} \right\},$$

may be replaced by

$$\frac{2(2s+1)}{s-1} \left( \frac{s+1}{\sqrt{\pi(s+\frac{1}{2})}} - 1 \right)$$

approximately, or by  $4(\sqrt{s/\pi} - 1)$ . Thus with  $s = 314$ , for a sample of 629 observations, the loss of information is near to 36 units, or the value of about 36 observations.

It is a matter of no great practical urgency, but of some theoretical importance, to consider the process of interpretation by which this loss can be recovered. Evidently, the simple and convenient method of relying on a single estimate will have to be abandoned. The loss of information has been traced to the fact that samples yielding the same estimate will have likelihood functions of different forms, and will therefore supply different amounts of information. When these functions are differentiable successive portions of the loss may be recovered by using as ancillary statistics, in addition to the maximum likelihood estimate, the second and higher differential coefficients at the maximum. In general we can only hope to recover the total loss, by taking into account the entire course of the likelihood function.

In our particular problem the curve of likelihood is a succession of exponential arcs, having  $n$  discontinuities at the values of the  $n$  observations of the sample, the exponent changing by  $-2$ , as each observation is passed in a positive direction. For the same value of our estimate, the median observation, this function will have very different forms according to the length of the intervals which separate the median from its successive neighbours. Any samples, however, in which these  $n-1$  intervals are the same will have the same

likelihood function. More explicitly the likelihood of the parameter having a value  $\phi$  as judged from the series of observations  $y_1, \dots, y_n$  will be equal to the likelihood of its value being  $\theta$  as judged from the series  $x_1, \dots, x_n$ , if

$$x - y = \theta - \phi$$

for each pair of observations in the pair of samples

We may specify the configuration of a sample by a series of positive non-decreasing numbers  $a_1, \dots, a_s$  representing the positive deviations from the median of the  $s$  largest observations, and a second series of positive non-decreasing numbers  $a'_1, \dots, a'_s$  representing the excesses of the median over the  $s$  smallest observations, so that if  $T$  is the median value the  $n$  observations are represented by  $T - a'_s, \dots, T - a'_1, T, T + a_1, \dots, T + a_s$ .

The probability of occurrence of any series of observations, the true centre of the distribution being  $\theta$ ,

$$L dx_1, \dots, dx_n$$

may now be written

$$n! L \frac{\partial (x_1, \dots, x_n)}{\partial (T, a_1, \dots, a_s, a'_1, \dots, a'_s)} dT da_1 \dots da_s da'_1 \dots da'_s \\ = n! L dT da_1 \dots da_s da'_1 \dots da'_s,$$

where, if, for example,  $\theta$  lies between  $T - a'_s$  and  $T - a'_{s-1}$

$$L = \frac{1}{2^n} e^{-(2s-1)(T-\theta) - S_1^2(s+s) + 2(s_1 + \dots + s_{s-1})}$$

Given the configuration of the sample, therefore, the probability that  $T$  lies in a range  $dT$ , between the limits  $\theta + a'_{s-1}$  and  $\theta + a'_s$ , is

$$df = \frac{1}{A} e^{2(s_1 + \dots + s_{s-1})} e^{-(2s-1)(T-\theta)} dT$$

of which the integral between these limits is

$$\frac{1}{(2s-1)A} e^{2(s_1 + \dots + s_{s-1})} (e^{-(2s-1)s_{s-1}} - e^{-(2s-1)s_s}),$$

and  $A$  is equal to the sum of all such integrals

$$1 - e^{-s'_1} + \frac{1}{2} (e^{-s'_1} - e^{2s_1 - 3s'_1}) + \frac{1}{2} (e^{2s_1 - 3s'_1} - e^{2s_1 + 2s_2 - 5s'_1}) + \\ + 1 - e^{-s'_1} + \frac{1}{2} (e^{-s'_1} - e^{2s_1 - 3s'_1}) + \frac{1}{2} (e^{2s_1 - 3s'_1} - e^{2s_1 + 2s_2 - 5s'_1}) +$$

Apart from the details of the analysis, however, it is apparent that *if attention is confined to samples having a given configuration* the sampling distribution of

$T$  for a given  $\theta$  is found from the likelihood of  $\theta$  for a given  $T$ , the probability curve in the first case being the mirror image of the likelihood curve in the second

To evaluate the amount of information supplied by this distribution we must evaluate the mean square of

$$\frac{d}{d\theta} \log \frac{df}{dT}$$

Now, if  $T$  lies between  $\theta + a'_{p-1}$  and  $\theta + a'_p$ ,

$$\log \frac{df}{dT} = - (2p - 1) (T - \theta)$$

so that in this case

$$\left( \frac{d}{d\theta} \log \frac{df}{dT} \right)^2 = (2p - 1)^2,$$

and the amount of information supplied by our estimate, in conjunction with a specification of the configuration of the sample from which it was obtained, is

$$\frac{1}{A} \{ 1 - e^{-a_1} + 3(e^{-a_1} - e^{2a_1 - 2a_1}) + 5( \\ + 1 - e^{-a_1} + 3(e^{-a_1} - e^{2a_1 - 2a_1}) + 5( \quad ) \}$$

This value will differ greatly from sample to sample. Thus, if  $a_1$  and  $a'_1$  were both large, so that the median lies in a considerable range otherwise unoccupied by observations, the amount of information approaches unity, at the other extreme if  $a_1$  and  $a'_1$  were both so small that  $e^{-a_1}$  is near to unity, then

$$A \rightarrow 2/(2s + 1),$$

and the amount of information rises to  $(2s + 1)^2$ , or  $n^2$

To find the average value of the amount of information derivable from the median, in conjunction with the configuration of the sample, we may note that the probability for a given configuration that  $s + p$  observations shall exceed, and  $s - p + 1$  fall short of the true value is

$$\frac{1}{(2p - 1) A} (e^{2a_1 + 2a_1} - (2p - 1) e^{a_1} - e^{2a_1 + 2a_1} - (2p - 1) e^{a_1})$$

and that the amount of information is obtained by multiplying this probability by  $(2p - 1)^2$  and adding for all values of  $p$

The average information for all configurations may, therefore, be found from the total probability for all configurations that exactly  $s + p$  observations

shall exceed the true value, since the probability of exceeding  $\theta$  is  $\frac{1}{2}$  independently for each observation, the probability is

$$\frac{1}{2^n} \frac{n!}{(s+p)! (s-p+1)!}$$

and this, multiplied by  $(2p-1)^2$ , and added for all values of  $p$ , will give the average amount of information. The probabilities are the terms of the expansion of

$$\left(\frac{1}{2} + \frac{1}{2}\right)^n,$$

and  $(2p-1)$  is twice the deviation from the mean corresponding to each value of  $p$ . The variance of the binomial is well known to be exactly  $\frac{1}{4}n$ , and the average amount of information used is consequently found to be exactly  $n$ , equal to the total amount known to be contained on the average in the sample.

The process of taking account of the distribution of our estimate in samples of the particular configuration observed has therefore recovered the whole of the information available. This process will seldom be so convenient as the use of an estimate by itself, without reference to the configuration, for instead of replacing the  $n$  observations by a single value, we now have to take account of all their values individually. Actually, indeed, in this case only the central group of values matters greatly, but in general the theoretical process illustrated here uses the available information exhaustively, only at the expense of abandoning the convenience of disregarding all properties of the sample beyond the best estimate it can provide. The reduction of the data is sacrificed to its complete interpretation.

The frequency distribution, which makes this complete interpretation possible, is the mirror image of the likelihood function. Thus if  $T_1$  is the estimate (the median) derived from the actual sample observed, and  $L(\theta - T_1)$  is the likelihood derived from this sample of any value of  $\theta$ , then the sampling distribution of  $T$  for any value of  $\theta$ , in samples of the same configuration is given by

$$df \propto L(\theta - T) dT$$

This is an extremely simple derivation of the sampling distribution of the estimate of maximum likelihood from the form of the likelihood function.

#### 4. The Simultaneous Estimation of Location and Scaling

In a very frequent class of cases not only the origin but the scale of the distribution is also represented by a parameter to be estimated from the observations. The frequency element is then of the form

$$f(\xi) d\xi,$$

where

$$\xi = \frac{x - \theta_1}{\theta_2}$$

In such cases it is obvious that the sample of values  $\xi$  in relation to any values  $\alpha_1$  and  $\alpha_2$  of the parameters corresponds in the sense of section 3 to the sample of values of  $x$  in relation to the values  $\theta_1 + \alpha_1\theta_2$  and  $\theta_2\alpha_2$ , and a double series of samples exists corresponding to any sample observed

The samples will have all the same configurations in the sense that supposing any two observations of the sample, such as the lowest and the lowest but one in value, have values  $a, a + b$  then the other members of the sample will be

$$a + bt_p, \quad p = 1, \dots, n-2,$$

where the  $n-2$  values of  $t_p$  specify the configuration, and are the same for all samples of which the configuration is the same

The frequency element

$$L dx_1, \dots, dx_n,$$

giving the frequency with which the  $n$  observations fall within assigned values, may then be replaced by

$$L \frac{\partial(x_1, \dots, x_n)}{\partial(a, b, t_1, \dots, t_{n-2})} da db dt_1 \dots dt_{n-2},$$

where the Jacobian is simply

$$\begin{vmatrix} 1 & 0 & 0 & 0 \\ 1 & 1 & 0 & 0 \\ 1 & t_1 & b & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & t_{n-2} & 0 & b \end{vmatrix}$$

or  $b^{n-2}$ . The simultaneous frequency distribution of  $a$  and  $b$  is therefore given by

$$df \propto L b^{n-2} da db$$

Now, it is evident that the estimates of  $\theta_1$  and  $\theta_2$  from such a sample will be

$$T_1 = a + \lambda b,$$

$$T_2 = \mu b,$$

where  $\lambda$  and  $\mu$  depend only on the configuration of the sample. Hence

$$\frac{\partial (T_1, T_2)}{\partial (a, b)} = \begin{vmatrix} 1 & 0 \\ \lambda & \mu \end{vmatrix} = \mu$$

and the distribution of these estimates in samples of the same configuration will be

$$df \propto LT_1^{n-2} dT_1 dT_2, \quad (4)$$

where in  $L$ ,  $T_1 + u_p T_2$  is substituted for  $x_p$ ,  $p = 1, \dots, n$ , the  $n$  values of  $u$  being known for the configuration observed.

If, therefore, we choose to take into account not merely the sampling distribution of our estimates for samples of all configurations, distributions which will involve, apart from the parameters of the population, only these two statistics, but rather the special simultaneous distribution for the particular configuration observed, we may obtain this special distribution directly from the form of the likelihood function.

Since, moreover, the whole course of the likelihood function is taken into account, it is, from this point of view evident that no information can be lost. An independent analytical proof of this is as follows, it is equally applicable to information in respect of  $\theta_1$  and of  $\theta_2$ .

The information respecting  $\theta_1$  contained in a single observation from the distribution (4) is numerically equal to the average value of

$$\left( \frac{\partial}{\partial \theta_1} \log L \right)^2$$

for all values of  $(T_1 - \theta_1)$  from  $-\infty$  to  $\infty$ , or, otherwise, to the average value of

$$\left\{ \frac{\partial}{\partial \theta_1} S(\log f) \right\}^2$$

where  $f(x - \theta_1)$  is the frequency of an observation falling in the range  $dx$ . The average for all values of  $T_1$  is, for any particular observation, the average for all values of  $x$ . Now the average value of

$$\frac{\partial}{\partial \theta_1} \log f$$

is zero, for

$$\int_{-\infty}^{\infty} \frac{\partial}{\partial \theta_1} \log f \cdot f \, dx = \int_{-\infty}^{\infty} \frac{\partial f}{\partial \theta_1} \, dx$$

which is zero, since the total frequency is unity, independent of  $\theta_1$ . But the average value for all values of  $(T_1 - \theta_1)$  and for all configurations including

variations of  $T_1$ , is the average value for all possible samples. We may apply this principle to the expression

$$S^2 \left( \frac{\partial}{\partial \theta_1} \log f \right)$$

when all the values of  $x$  are independent. Then the average value of the square of the sums of  $n$  terms, independent and all having a mean value zero, is  $n$  times the mean square of each of them, or  $n$  times the mean value of

$$\left( \frac{\partial}{\partial \theta_1} \log f \right)^2$$

for all values of  $x$  from  $-\infty$  to  $\infty$ , which is, by definition, the amount of information supplied by a sample of  $n$  observations. Hence the average amount of information respecting  $\theta_1$  supplied by (4) for all configurations is the entirety of that supplied by the data.

With respect to  $\theta_2$ , we require the average value of

$$S^2 \left( \frac{\partial}{\partial \theta_2} \log f \right)$$

for all values of  $T_2$  from 0 to  $\infty$ . The average of this for all configurations and for all values of  $T_1$ , again reduces to the mean value of

$$n \left( \frac{\partial}{\partial \theta_2} \log f \right)^2$$

for all values of  $x$  from  $-\infty$  to  $\infty$ , and so to the average amount of information contained in a sample of  $n$  observations.

### Summary

(I) Reasons are given for the use of mathematical likelihood in problems of inductive inference.

(II) When a statistic exists, satisfying the criterion of sufficiency, the likelihood function involves only that statistic.

(III) An example is given of a sufficient statistic, and its sampling distribution is expressed in terms of the likelihood function.

(IV) This property is generalized for all cases of simple estimation, where a sufficient statistic exists.

(V) It is shown that these cases and only these supply tests of significance of the kind termed by Neyman and Pearson "uniformly most powerful" with regard to a class of alternative hypotheses.



(VI) Where no sufficient statistic exists the precision of estimation may in general be enhanced by the use of ancillary statistics. A class of cases is defined and illustrated in which the totality of the ancillary information supplied by the observations may be utilized.

(VII) This process gives a very simple derivation of sampling distributions, in which there is no loss of information, even for small samples.

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*The Electrical Condition of Hot Surfaces during the Adsorption of Gases. Part V—The Charging up of Hot Surfaces*

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(Communicated by W. A. Bone, F.R.S. - Received July 31—Revised December 15, 1933)

*Introduction*

The investigation into the electrical condition of hot surfaces has consisted mainly of measurements of (a) the steady equilibrium potentials acquired by such surfaces when heated in a vacuum or in contact with various gases, and (b) the rates at which these potentials build up after earthing. The previous communications of this series\* have dealt with the steady potentials exhibited by gold, silver, nickel, platinum, carbon, and copper surfaces under varying conditions. During the course of these experiments the rates of attainment of the steady potentials were also observed and recorded, and are described below. The results with the silver sheet, however, have been omitted for reasons which have already been given. Certain general regularities in the results have been distinguished, whereby much light is thrown on the nature and mechanism of the processes taking place at the interface between solid and gas phases. It has now been found possible to advance a more complete explanation of the changes occurring on the surfaces and which give rise to their electrical charging up as observed in these researches. It also gives information, which is of fundamental importance, on the subject of the role played by hot surfaces in heterogeneous catalytic reactions.

\* 'Proc. Roy. Soc.,' A, vol. 116, p. 379 (1927), vol. 120, p. 235 (1928), vol. 124, p. 356 (1929), and vol. 132, p. 192 (1931).

*Experimental*

The apparatus employed has been described for the most part in sufficient detail in the earlier communications. The addition of a number of multi-vane variable air condensers, each of 0.001 microfarad capacity in parallel to the surface-electrometer-earth system made it possible to vary the capacity of any surface to earth within suitable limits. The condensers, connecting leads, and switches were completely enclosed in an earthed metallic shielding as a protection against spurious electrical effects from external sources. The

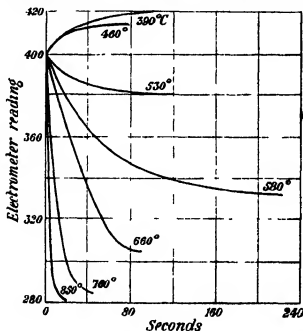


FIG 1

condensers were calibrated against a Dubilier standard condenser, while the capacities of the surface-electrometer systems were determined by the method of mixtures, one of the variable air condensers serving as a sub-standard.

*The Rate of Charging Up*—The rates of charging up of the surfaces were calculated from the observed electrometer readings at suitable time intervals starting from zero potential, *i. e.*, at the moment of re-insulation of the surface-electrometer system after earthing. The general type of the curves connecting the readings of the electrometer with time is shown in fig 1. These were obtained in the course of experiments with the nickel sheet in contact with oxygen at a constant capacity of 1020 cm (Part II, Series C, and the present paper, Series B), and they represent the general characteristics of the manner in

which the hot surfaces charge up to the equilibrium potentials. The slope of such curves gives a measure of the current ( $i$ ) flowing from the surface at any instant, and can be calculated by the equation

$$i = C \, dv/dt, \quad (1)$$

where  $C$  is the capacity of the surface-electrometer system,  $dt$  is the interval of time between two electrometer readings, and  $dv$  is the corresponding incre-

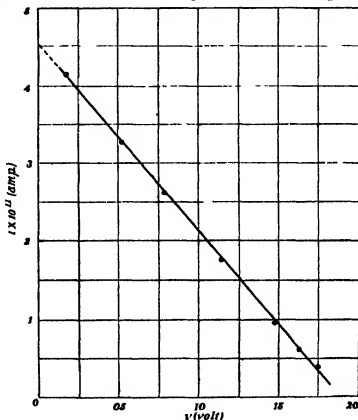


FIG. 2

ment of potential. The current thus obtained was taken to correspond to the potential at the middle of the time interval. This method of calculation was also used by Richardson and Brown\* in their experiments on the kinetic energy of electrons and positive ions emitted from hot bodies by measuring the rate of charging up of a neighbouring electrode. Values calculated in this way using equation (1) for the nickel surface in contact with oxygen at 580° C, from the charging up curve given in fig. 1, have been plotted in fig. 2.

\* 'Phil Mag.' vol 16, p. 353 (1908), vol 17, p. 355 (1909), and vol 18, p. 649 (1909).

The relationship between the momentary current flowing from the surface ( $i$ ) and the potential of the surface at the same time ( $v$ ) is very nearly linear, and can be represented by an equation of the general type

$$i = i_0 - av, \quad (2)$$

where  $i_0$  is the value of  $i$  when  $v = 0$ ,  $i_0$ , the instantaneous value of the current flowing from the surface at the moment of insulation from earth, and  $a$  is a constant equal to the slope of the straight line. These calculations have been applied to a large number of the experimental results and the approximately linear relation between the current from the surface at any time during the charging up process and the mean potential of the surface during the same time has always been obtained. Richardson and Brown also found that there was a tendency for the current to be a linear function of the potential, especially after a long period of heating in a vacuum, or at a high temperature. The current flowing from the surface is zero when its potential has reached the limiting value,  $V$ , under the given experimental conditions.

The rates of charging up in Series A to E recorded below have been expressed as values of  $i_0$ ,  $i_0$ , the instantaneous maximum current flowing from a surface just as it is insulated from earth. Two methods were employed in this investigation for the evaluation of  $i_0$ . (a) the straight line obtained by plotting  $i$  against  $v$  in the manner outlined above was extrapolated to the axis of  $i$  to give the value of  $i_0$ , (b) it could also be calculated from  $I$  the average or mean rate of charging up to a potential equal to half the limiting value ( $V/2$ ). It can be shown that

$$I = CV/2t = \frac{1}{2}i_0,$$

where  $t$  is the time in seconds required to attain half the limiting potential, in volts, in an electrometer system of capacity  $C$ , in farads. The values of  $i_0$  are independent of the actual capacity of the surface-electrometer systems as the rate of increase of potential on the surface was always found to vary inversely with capacity.

As the rate of charging up of a surface in any given experiment can be expressed in the form of an equation of the first degree, it follows that it has been completely and fully represented in terms of the equilibrium potential,  $V$ , and the maximum rate of charging up,  $i_0$ .

Equation (2) can be rewritten in the form

$$C \, dv/dt = a(V - v),$$

where  $C$  is the capacity of the surface-electrometer system,  $V$  is the final

steady value of the surface potential, and  $v$  is the momentary potential at any time  $t$ . By integrating and re-arranging terms this becomes

$$v = V(1 - e^{-at/C})$$

The fact that the variation with time of the potential acquired by a surface could be represented by a function of this form has also been indicated by independent work in these laboratories \*

### *Experimental Results*

It has been repeatedly confirmed throughout the course of this investigation that the magnitude of the steady surface potential in contact with a gas was independent, within the limits of experimental error, of the gaseous pressure between 100 mm and atmospheric pressure. In a similar way it has been found that within this range the rate of charging up of a surface to the steady potential characteristic of the given experimental conditions does not depend on the pressure. The results given below have been confined to experiments with single gases, as those with reacting mixtures require more detailed examination especially in connection with their rates of interaction.

*Series A Experiments with the Gold Sheet*—The surface potential results of this series of experiments have been previously described in Part I (Series C). The rates of charging up, as expressed by  $\tau_0$ , for this surface are recorded against temperature up to 850° C (in *vacuo* and in contact with hydrogen, oxygen, argon, carbonic oxide, and nitrogen) in fig. 3. The dotted portion of the oxygen curve is that over which the potential was changing from a positive to a negative value and where it is very difficult to obtain reproducible results.

On admitting a gas into contact with the heated gold sheet when it was exhibiting the appropriate *in vacuo* surface potential, it has been shown that the latter gradually gave place to the potential characteristic of the particular gas during a period of time which was dependent on the nature of the gas. It has also been found that this change is accompanied by a gradual and similar alteration in the rate of charging up, while removal of a gas by evacuation produces the same type of change in the reverse direction. This behaviour has been confirmed in the experiments with the other surfaces, the appropriate change in the rate of charging up always accompanying such an alteration in the experimental conditions.

*Series B Experiments with the Nickel Sheet*—The variations in the rates of charging up of the nickel sheet with temperature, corresponding to the surface

\* Bradford, "Dissertation, Ph D.," London (1931)

potentials determined in Part II (Series C, D, and F), are given in fig. 4. The curves recorded are for hydrogen, *in vacuo*, carbonic oxide, oxygen, argon, and nitrogen.

*Series C Experiments with the Platinum Sheet*—The results correspond to the surface potential experiments described in Part III (Series A, B, and C). In the experiments with the platinum sheet heated only to a maximum tempera-

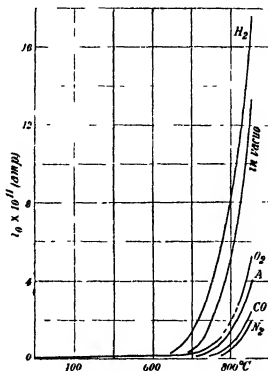


FIG. 3.

ture of 500° C the values of  $i_0$  were very small, even at 500°, thus hydrogen (1),  $0.04 \times 10^{-11}$  amp, *in vacuo*,  $0.33 \times 10^{-11}$  amp, and hydrogen (2),  $0.04 \times 10^{-11}$  amp. In the series up to 660° the rates of charging up at 660° were *in vacuo* (1),  $0.79 \times 10^{-11}$  amp, hydrogen,  $0.80 \times 10^{-11}$  amp, oxygen,  $0.76 \times 10^{-11}$  amp, and *in vacuo* (2),  $2.00 \times 10^{-11}$  amp. Similar measurements were carried out during the series of experiments up to 850° and are recorded in fig. 5.

*Series D Experiments with the Carbon Rod*—The rates of charging up of the carbon rod corresponding to the potential measurements recorded in Part IV (Series A) are plotted against temperature in fig. 6. Most of the values

of  $i_0$  have been given only over the lower part of the temperature range (up to  $90 \times 10^{-11}$  amp), as at higher temperatures the increases were so very

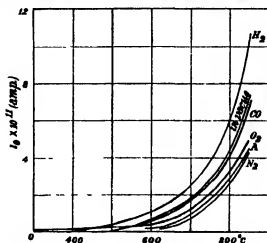


FIG 4

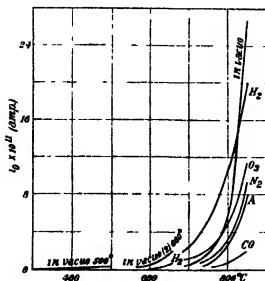


FIG 5

rapid that  $i_0$  was, for the most part, beyond the limits of accurate experimental measurement.

*Series E. Experiments with the Copper Sheet.*—The rates of charging up in the following experiments, already described in Part IV (Series B), with the

copper sheet are recorded in fig 7 *in vacuo* and hydrogen up to 520°, *in vacuo* and hydrogen up to 660°, and *in vacuo*, hydrogen, nitrogen, and carbonic oxide up to 850°

*Series F Experiments on the Rate of Leak of a Potential applied to a Hot Surface*—A large number of experiments have been carried out with the surfaces so far examined under various conditions of temperature and gas pressure, in which the rate of leak of an applied potential was determined. The

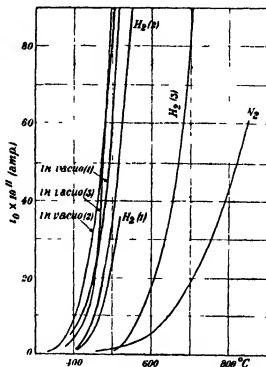


FIG 6

experimental procedure was as follows. The potential of the surface-electrometer system was brought to a known positive or negative value whereupon it was left insulated while electrometer readings at suitable time intervals were observed. It was found that the potential of the system always changed in a gradual and uniform manner, from the applied value to that characteristic of the surface under the given experimental conditions. The rates at which applied voltages of + or - 1, 2, and 4 volts leaked away from a heated gold gauze in contact with oxygen, recorded in fig 8, illustrate the general nature of these changes.



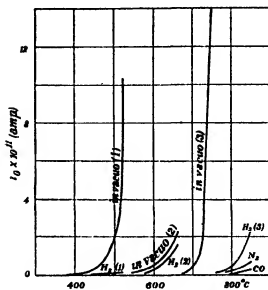


FIG 7

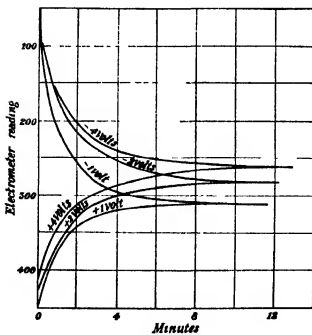


FIG 8

Bangham and Lewis\* have investigated the effect of gases on the electrical charges developed by heated metals, and have found, in experiments similar to those described in this series, that the mode of approach to the equilibrium potential was widely different according to whether equilibrium value was approached from the positive or from the negative side. In the latter method, they found that the rate of change was a highly sensitive function of the momentary potential, in the former it was markedly insensitive, the rate being nearly uniform. It has not been found possible, during the course of the present investigation, to reproduce Bangham and Lewis' results, in fact, smooth and regular changes of the type shown in fig. 8 have been obtained in every experiment so far carried out. It might be suggested that Bangham and Lewis had not been entirely successful in overcoming all the experimental difficulties to be met with in measurements of this nature.

#### *Discussion of the Results*

The most important facts revealed by a study of the rates of charging up of the various surfaces are as follows —

(a) When the rates of charging up, expressed as values of  $\tau_0$ , for the gold, nickel, and platinum surfaces at 850° C are tabulated as in Table I, it is seen that they fall naturally into a group exhibiting certain well-defined features

Table I — (Multiplied by  $10^{11}$ )

Surface	H <sub>2</sub>	in vacuo	O <sub>2</sub>	A	CO	N <sub>2</sub>
Platinum	20.00	26.67	11.27	8.00	2.00	9.33
Gold	17.60	13.33	5.33	4.03	2.53	2.00
Nickel	10.60	7.36	5.00	4.53	7.07	4.32

This shows that the rates of charging up for the platinum sheet were larger throughout than those for gold or nickel except in contact with carbonic oxide where the order was completely reversed. It is quite reasonable to suspect that the low rate for platinum under these conditions is probably connected with the well-known poisoning effect of carbonic oxide on platinum catalysts,† whilst the higher rate for nickel in contact with carbonic oxide is associated with the method of attachment of these molecules to the surface as indicated in Part II. For the gold and nickel sheets, beyond the fact that the rates in

\* *J. Chem. Soc.*, p. 1140 (1929)

† Rideal and Taylor, "Catalysis in Theory and Practice," p. 133 (1936).

contact with hydrogen and *in vacuo* always exceeded those in contact with other gases, no general regularity seems to exist

(b) In the experiments with the carbon rod the rates of charging up were very large compared with those considered in (a). Thus, with the carbon rod *in vacuo* (3)  $i_0 = 93.3 \times 10^{-11}$  amp at  $500^\circ$  and was rising extremely rapidly, in contact with hydrogen (3)  $i_0 = 86.7 \times 10^{-11}$  amp at  $700^\circ$  and was likewise rising rapidly, in contact with nitrogen  $i_0 = 62.0 \times 10^{-11}$  amp at  $850^\circ$ . Even this latter value was nearly seven times greater than the corresponding rate for the platinum sheet. The apparent area of the surface of the carbon rod was almost of the same order as that of the metal sheets, but the work of Bowden and Rideal\* has shown that its accessible area, *i.e.*, that which could be reached by a reactant, might be over 300 times its apparent area. Furthermore, there are a number of substances present in the type of carbon rod used in this investigation which might have had the effect of increasing the rates of charging up. Thus it seems quite likely that the rapid rates of charging up of the carbon rod were due, at least in part, to one or other or both of these causes. It has been pointed out in Part IV that previous heat treatment had a profound effect on the values of the surface potentials of the carbon rod *in vacuo*, whereas the rates of charging up seemed to depend very little on heat treatment. Thus, in fig. 6 the *in vacuo* curves (1), (2), and (3) were very close over the whole temperature range where accurate measurements were at all possible. Similarly, the rates of charging up for the carbon rod in hydrogen heated to maximum temperatures of  $520^\circ$  and  $660^\circ$  respectively were very nearly identical. Heat treatment at  $850^\circ$ , however, greatly influenced the results, *e.g.*, at  $550^\circ$  the rate of charging up was reduced from 93.3 to  $6.7 \times 10^{-11}$  amp.

(c) The rate of charging up of the copper sheet *in vacuo* at temperatures above  $750^\circ$  was extremely high, but in contact with gases, including hydrogen, it was low compared with the other metals. At the present time it is not possible to advance any explanation of this behaviour, but it is quite probable that on completion of an investigation now in progress in these laboratories further light may be thrown on this subject.

(d) In Part IV it was established that the steady potentials exhibited by surfaces under similar experimental conditions tend to approach to zero as the normalization temperature is raised. The results with the platinum and copper sheets in Series C and E indicate that this change is accompanied by a reduction in the rate of charging up, and this can be illustrated by the figures given in Table II.

\* 'Proc Roy Soc,' A, vol 120, p. 80 (1928)

Table II

Surface	Temperature of normalization	Experimental temperature	( $t_0 \times 10^{12}$ ).
Platinum in vacuo	° C 500 660	° C 500 500	amp. 0 33 0 07
	660 850	660 660	2 00 0 27
	520 660	520 520	10 33 0 13
	660 850	660 660	2 24 0 05
Copper in vacuo	520 660	520 520	10 33 0 13
	660 850	660 660	2 24 0 05
	520 660	520 520	10 33 0 13
	660 850	660 660	2 24 0 05

(c) Figs. 3 to 7 show that the rates of charging up of hot surfaces increase rapidly with temperature. It has been found that the logarithm of the rate of charging up of a surface is a linear function of the reciprocal of its absolute temperature. This relationship can be expressed by the equation

$$\log i_0 = \log a - b/T, \quad (3)$$

where  $a$  and  $b$  are constants. It follows that

$$i_0 = ae^{-b/T},$$

$e$  being the base of the natural logarithms. Fig. 9 gives the results of plotting values of  $\log(i_0 \times 10^{12})$  as ordinates against values of  $10^3/T$  as abscissae for the carbon, nickel, platinum, gold, and copper surfaces respectively in contact with hydrogen, and for each  $\log i_0$  is seen to be a linear function of  $1/T$ . The special significance of equation (3) can be seen when it is compared with the Arrhenius equation for the temperature variation of the velocity constant,  $k$ , for any reaction

$$\log k = C - A/RT \quad (4)$$

My thanks are due to the Department of Scientific and Industrial Research for a grant which enabled me to devote my full time to this research, and to Professor G. I. Finch for his continued interest and assistance.

### General Summary

The rates at which the steady equilibrium potentials are built up on gold, nickel, platinum, carbon, and copper surfaces after earthing have been studied

under varying experimental conditions. The experiments have shown that the rate of charging up of a surface is a linear function of its instantaneous potential. The effect of the surface temperature is very marked, the logarithm of the rate of charging up being a linear function of the reciprocal of the

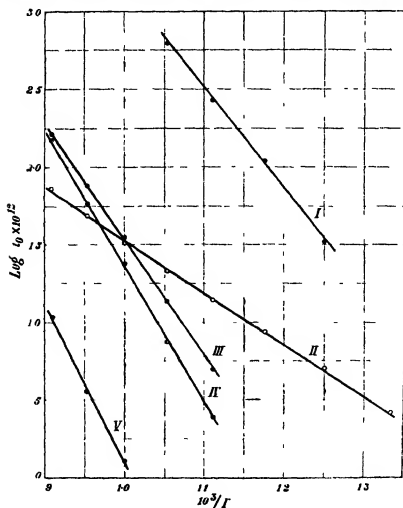


FIG 9

absolute temperature. According to the evidence available it is extremely probable that the hot surfaces emit positive electricity over the temperature range investigated (up to  $850^\circ \text{C}$ ). When heated in a vacuum there is little doubt that the emission consists of positively charged metal ions, while in experiments carried out in contact with gases, the ions are positively charged

atoms or molecules of the gas. With oxygen at low temperatures, however, the ions appear to be negatively charged. The ionic emission is preceded by a partial separation of the atoms of the gas molecules by adsorption with the formation of heteropolar molecules. When the free atom of one of the latter can acquire sufficient thermal energy, it is liberated as an ion, the sign of which depends on the orientation of the polarized molecule on the surface.

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*The Electrical Condition of Hot Surfaces Part VI — A Gold Surface  
Catalysing the Combustion of Carbonic Oxide*

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Technology

(Communicated by Professor W. A. Bone, F.R.S. — Received July 31 — Revised  
December 15, 1933)

*Introduction*

A study of the electrical condition of hot surfaces in contact with gases\* has revealed the fact that in several important respects a striking parallel exists between such electrical condition and the catalytic properties of the surfaces. Thus, it has been found that, whereas at comparatively low temperatures the specific effect of a surface upon the potential it acquires in contact with a gas is highly pronounced, with increasing temperature such specificity tends to disappear until, at a sufficiently high temperature, the value of the surface potential is almost wholly determined by the nature of the gas and is nearly independent of the nature of the surface itself. This may be compared with the fact, established by Bone and his co-workers,† that the differences between the catalytic powers of various surfaces, which are often considerable at low temperatures, diminish as the temperature is raised, until they practically disappear. Furthermore, it is now well recognized that the catalytic properties of a surface are intimately connected with its structure, and a close parallelism has been observed between the surface potential of a metal and changes in its

\* See footnote, p. 307

† 'Howard Lectures, Royal Society of Arts,' 1914

superficial structure brought about by heat treatment.\* Finally, it has been found that the electric current which can be drawn from a heated metallic surface in contact with a gas increases rapidly with increasing temperature,\* an observation which has its parallel in the well-known rapid increase in the catalytic activity of such a surface with increasing temperature.

The present investigation was undertaken with the object of determining the relationship, if any, between the role played by moisture in promoting the heterogeneous catalytic combustion of carbonic oxide with oxygen and the structure of the surface as revealed by its catalytic activity and electrical condition.

### Experimental

In these experiments, moist or dry mixtures of carbonic oxide and oxygen in their combining proportions were circulated over an electrically insulated, heated gold gauze surface connected to a Landemann quadrant electrometer. The reaction velocity, the rate of charging up and the equilibrium potential of the surface were recorded. We have found, *inter alia*, that (i) the structure of the surface most suitable for the promotion of the combustion of dry carbonic oxide differed from that required when steam was present, (ii) the reaction velocity was greater in the presence of steam than otherwise, with increasing temperature, however, the rates of reaction in the moist and dry systems converged until in the neighbourhood of  $550^{\circ}\text{C}$  they were nearly equal, and, finally, (iii) changes in the activity of the surface were paralleled by changes in the rate of electrical charging.

*The Apparatus*—The combustion of  $2\text{CO} + \text{O}_2$  mixtures in contact with a heated gold gauze was carried out in one or other of the two circulation systems shown in fig. 1, according to whether the moist or the dry reaction was being studied. The gold gauze, G, was suspended on a quartz rod in the electrically heated transparent quartz vessel, S, which was joined into the circulation systems by graded seals at *s* and *s'*. A gold wire welded to the gauze led out of the combustion vessel through a quartz side tube, sealed off with sealing wax, to a Landemann electrometer. The  $\text{CO}_2$  absorption and drying tubes were each 5 feet long. Fig. 1 is self-explanatory so far as the remainder of the apparatus is concerned. All parts of the apparatus were cleaned before assembly with hot nitric-chromic acid cleaning mixture, and then steamed out with ammonia-free steam. The individual parts of the dry system were further dried and strongly heated while passing a current of dry oxygen. Precautions were

\* *Loc. cit.*, Part I, p. 395, and Part V, p. 367.

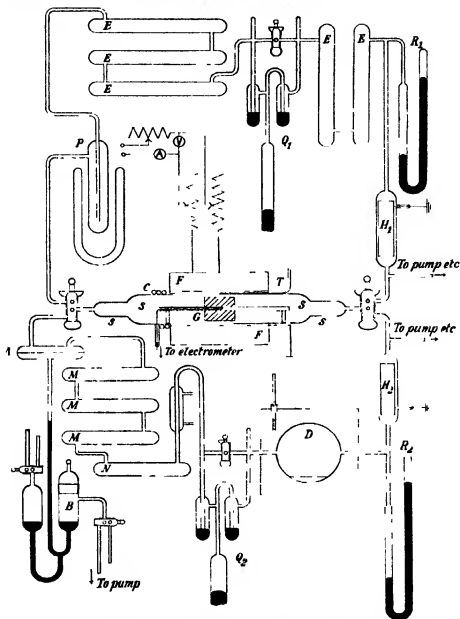


FIG 1—SS—combustion vessel ending in graded quartz to glass seals. G—gold gauze C—water-cooling coil T—thermocouple F—electric furnace, non inductively wound with nichrome over an iron tube which, together with the electrical mid point of the winding, was carthed  $Q_1, Q_2$ —dry and moist system circulating pumps respectively  $H_1, H_2$ —earthed aluminum foils  $M, M, M$ —moist system absorption tubes, packed with 1:10 soda lime N—guard tube parts ally filled with saturated baryta solution A—short absorption tube containing concentrated sulphuric acid to dry gas roughly before removal of  $\text{CO}_2$  B—a device enabling the spent acid to be replaced without allowing air to enter the apparatus. D—1-litre bulb containing distilled water and surrounded by heat-insulating material, to maintain constant water-vapour pressure in the reaction zone. F—liquid-air cooled trap for removal of  $\text{CO}_2$  in dry system  $E, E$ , etc—drying tubes filled with redistilled phosphoric oxide.  $R_1, R_2$ —closed-limb mercury manometers.



taken to prevent the ingress of moisture to the dry system during assembly. The purified tap-grease recommended by Baker\* was used as lubricant throughout the apparatus, the grease was thoroughly out-gassed before use.

*The Circulating Pumps* †—In all the experiments to be described below the rate of circulation was such that the volume of gas, measured at the temperature and pressure of the apparatus, circulated in unit time was constant. The time of one complete circuit, whether in the moist or dry system, was 13 minutes. This uniform rate of circulation was obtained by maintaining a constant oscillation frequency of the mercury piston and controlling its amplitude by suitable weighting of the plunger.

*The Gases*—Carbonic oxide and oxygen were prepared and purified in the manner previously outlined ‡. The purity of the gases was verified in two ways: (i) by analysis in a Bone-Newitt apparatus, and (ii) by determining both the insensitiveness to ignition by a condensed discharge and the dielectric strength§ of a rigidly dried equivalent mixture of the two gases, a procedure which afforded a stringent test for the absence of hydrogen and hydrogenous gases. The dry system was filled with the equivalent mixture through three tubes 1 in. long, containing redistilled phosphoric oxide.

*The Gold Gauze*—The history and other details of the gold gauze have been given previously ||. Since completion of Finch and Stimson's experiments, in the course of which the gauze was last used, it had been stored for a period of three years in dry air.

*Experimental Procedure*—The rate of catalytic reaction was determined first in moist, then in dry, and finally again in moist  $2(\text{CO}) + \text{O}_2$  experiments being carried out in each series at several temperatures. The gauze was never allowed to cool to room temperature during the course of any of these three main series of experiments and was kept as far as practicable in contact with the circulating mixture, long periods of stagnation or evacuation being avoided.

The range of pressures employed was from 350 mm. down to the lowest limit of effective circulation, approximately 10 mm. The time required for complete combustion in the case of either of the reaction systems over this pressure range varied from several weeks at the lowest to less than one hour at the highest temperatures. Manometer readings were corrected for temperature

\* 'J. Chem. Soc.', vol. 128, p. 1661 (1920).

† *Ibid.*, vol. 127, p. 2464 (1925).

‡ 'Proc. Roy. Soc. A', vol. 124, p. 303 (1929).

§ 'J. Chem. Soc.', vol. 129, p. 1540 (1930).

|| 'Proc. Roy. Soc. A', vol. 116, p. 379 (1927).

variations, standardized thermometers being placed at different parts of the apparatus for this purpose. An additional correction was made in experiments in the dry system to allow for variation in the level of the liquid air surrounding the carbon dioxide trap. In comparing rates of reaction at different temperatures, a small correction was necessary in order to take into account corresponding changes in the mass of gas contained in the heated zone of the furnace at different temperatures. Differences of volume had also to be allowed for in comparing rates of reaction in the moist and dry systems. The rates of reaction were derived from the mean slopes of the graphs connecting the logarithms of the corrected partial pressures of the dry nitrogen-free gas with time. Samples of the gas entering the reaction vessel were frequently taken and analysed in order to test the efficacy of the  $\text{CO}_2$  absorption tubes, and to verify the absence of leaks. In the moist system, tests for the presence of hydrogen were also applied, but this gas was never found.

The rate of charging-up of the surface on re-insulation after earthing was found to proceed in accordance with the equation  $v = A(1 - e^{-Pt})$ , where  $v$  is the surface potential at the time,  $t$ ,  $A$ , is the final equilibrium, and therefore maximum, value of the surface potential, and  $P$  is a constant for given conditions of temperature, gas pressure, and surface activity. Hence, the product  $PC$ , where  $C$  is the electrical capacity of the surface system, afforded a measure of the rate of charging-up of the surface. In what follows the values of  $PC$  are expressed in terms of micro-micro-coulombs per minute per volt.

### *The Results*

Fifteen series of experiments were carried out. In the first two, moist (15 mm  $\text{H}_2\text{O}$ )  $2\text{CO} + \text{O}_2$  mixtures were circulated over the gold gauze at  $245^\circ$  and  $334^\circ$  respectively, in the next five series, dry mixtures were employed, the temperature range being between  $245^\circ$  and  $495^\circ$ , finally, eight series were carried out in a moist system between  $228^\circ$  and  $530^\circ$ . Throughout, the gauze was never in contact with any gas other than dry or moist  $2\text{CO} + \text{O}_2$  mixtures.

A fully detailed account of all the experimental results would occupy much space and is hardly necessary in order to elucidate the more important facts. It will suffice to give in the first place a broad and general summary of the main experimental results, and then where necessary to set forth in sufficient further detail those results whereby such facts are established which cannot be brought out, or are insufficiently emphasized in the more general summary.

The results are set forth in summarized form in Table I and fig. 2

Table I

Condition	Series	Temperature °C	Time of heating	Initial stage			Intermediate stage			Final stage		PC
				I	A	PC	I	A	PC	k	A	
Moist	1	245	38 days	0 0062	-0 70	8 0	0 0076	-1 75	5 5	0 0077	- & -	-
	2	334	10 days	0 0330	+0 55	4 5	0 0305	+ & -	-	0 0285	+1 35	1 4
	3	245	18 days	0 0006	-0 14	0 4	0 0010	-0 15	0 3	0 0012	+0 20	0 3
Dry	4	325	7 days	0 0030	-0 07	1 6	0 0036	+0 09	0 5	0 0036	+0 10	0 5
	5	410	14 days	0 0230	-0 09	0 18	0 0200	-0 19	0 20	0 0260	-0 35	0 24
	6	495	4 days	0 0530	-0 82	38	0 0425	-0 86	36	0 0420	-0 86	34
	7	565	3 days	0 0680	-0 51	275	0 0570	-0 51	185	0 0615	-0 51	160
	8	226	1 day, 2 months cold, 14 days	First day			After 2 months			After further 14 days		
Moist	9	228	15 days	0 0036	+0 13	0 02	0 0026	-0 26	0 70	0 0026	+0 20	0 20
	10	317	12 days	0 0039	-3 8	1 3	0 0045	-0 10	2 0	0 0046	+0 50	1 0
	11	406	9 days	0 0120	+0 25	0 50	0 0112	-ve	-	0 0109	+ & -	-
	12	484	19 days	0 0280	-ve	-	0 0340	- & -	-	0 0350	-ve	-
	13	530	5 days	0 6070	-0 50	21	0 0592	-0 60	16	0 0379	-0 55	10
	14	470	2 days	0 160	-0 75	130	0 190	-0 75	215	0 122	-0 76	186
	15	373	2 days	0 0660	-0 48	10	-	-	-	0 0560	-0 49	17
	16	485	1 day	0 0225	-0 15	0 49	-	-	-	0 0200	-0 15	0 54
				0 0570	-0 45	5 1	-	-	-	-	-	-

Note—Values of  $k$ ,  $A$ , and  $PC$  above refer to a total gas pressure of 100 mm except in Series 3 and 9, in which the pressures were 260 mm and 160 mm, respectively

These results establish the following facts —

(1) On comparing the rates of reaction in the dry and moist systems it will be seen that (a) under similar experimental conditions, the reaction velocities were invariably higher in the moist than in the dry system. With increasing temperature, the difference between the rates observed in the two systems diminished practically throughout the range of temperature covered by the experiments. These observations confirm the results obtained by Bone\* for the rate of combustion of a  $P_2O_5$ -dried  $2CO + O_2$  mixture on the surface of the same

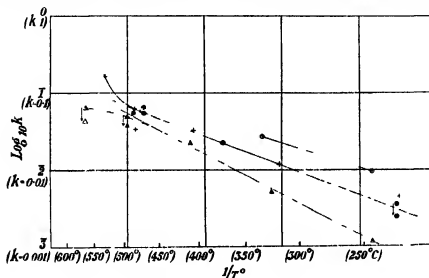


FIG. 2

Moist	$\left\{ \begin{array}{l} + \text{ Series 9 to 13} \\ \circ \text{ Series 1 and 2} \end{array} \right\}$	temperature raised throughout
	$\oplus$ Series 14 to 16,	temperature varied.
Dry	$\left\{ \begin{array}{l} \triangle \text{ Series 3 to 7, temperature raised throughout} \\ \bullet \text{ Series 8(a) and (b)} \end{array} \right\}$	

specimen of gold gauze at a temperature of  $240^\circ C$ , and give strong support to the suggestion made by that author that at sufficiently high temperatures the reaction of carbonic oxide and oxygen might prove to be independent of the presence of moisture. At the highest temperatures, however, the reaction velocity in the moist system attained what appeared to be an abnormally high value, whereas in the dry system,  $k$  showed a similarly abnormal decrease, (b) on changing from the moist to the dry reaction or *vice versa*, a long period of time elapsed before complete readjustment,  $\pm e$ , normalization of the surface to the

\* 'Proc Roy Soc,' A, vol 112, pp 474-499 (1926)

new conditions, was attained (see Series 3 and 9), (c) after drying, the efficiency of the surface in promoting the moist reaction was found to be permanently impaired (*cf.* Series 1 with 9, or 2 with 10, see also fig 2). The apparent heat of activation of the moist reaction was not, however, altered as a result of drying, and, furthermore, the reduction in the value of  $k$  at a given temperature was practically equal to the total reduction in the activity of the surface at high temperatures in the dry system.

(2) On maintaining the surface at constant temperature for long periods of time, the catalytic activity did not in general remain constant, thus, (a) when first heated to 245° in the moist system (Series 1), the reaction velocity increased rapidly at first, and then more slowly. Table II illustrates the rapid initial rise.

Table II

Time after attainment of constant temperature (hours)	0	0.75	1.42	1.83	2.33	1.17	4.08	4.07
Reaction velocity constant, $k$	0.00096	0.00235	0.00238	0.00308	0.00368	0.00397	0.00437	

After a further period of 18 hours the observed pressures and times gave a sensibly linear relationship between  $\log p$  and  $t$ , the value of  $k$  being 0.00630. The catalytic activity of the surface continued to increase slowly for 37 days, as shown in Table I (Series 1). (b) At higher temperatures in both moist and dry systems, other effects became evident. Thus, in the dry system over the range 300°–400°, the values of  $k$  increased slowly with time, whereas, at temperatures above 450°, the values of  $k$  at constant temperature fell from the commencement of each series. In the moist system, apart from the initial normalization referred to above, a slight decrease in the value of  $k$  was observed in Series 2 at 334° and in Series 10 at 317°. In Series 11 at 406° a gradual rise occurred similar to that at the same temperature in the dry system (Series 5). At 484° (Series 12),  $k$  fell steadily, but, as will be shown later, the fall could not be regarded as of the same nature as that which took place in the dry system. At the highest temperature attained in the moist system (Series 13 at 530°), the values of  $k$  passed through a maximum.

(3) Slow changes in the value of  $k$  at a given temperature were not necessarily accompanied by permanent changes in the catalytic activity of the surface at other temperatures or in different conditions. Thus, although the gradual decrease in  $k$  at the higher temperatures in the dry system led to an approximately proportional decrease in the catalytic activity in the moist, it also

resulted in increased activity at low temperatures in the dry (see Series 3 and 8). Also, during the experiments in the moist system at  $484^{\circ}$  (Series 12), the value of  $k$  decreased to 60% of its initial value. Further changes took place at  $530^{\circ}$  (Series 13), where, although  $k$  attained a maximum, its value at the close of the series was lower than at the commencement. In spite of these decreases, on returning to  $470^{\circ}$  (Series 14),  $k$  had practically the same value as at the commencement of Series 12, and at a lower temperature ( $373^{\circ}$ , Series 15) the value of  $k$  agreed well with that interpolated between Series 10 and 11, previous to the high temperature treatment. Hence it may be concluded that the gradual decrease in  $k$  in Series 12 did not entail a permanent decrease in the catalytic activity of the surface at other temperatures.

(4) With few exceptions, on changing from a lower to a higher temperature, the value of  $k$  showed a rapid initial increase and, for the reverse temperature change, a rapid initial decrease. These effects were, however, sometimes masked to some extent by the occurrence of other changes in the catalytic activity at constant temperature. The initial increase on passing from a lower to a higher temperature may be seen in Series 1, similar effects in Series 3, 4, 5, 9, and 11 may not have been due solely to temperature increase. In Series 13 an initial increase was followed by a decrease. The reverse effect is clearly shown in Series 8. Thus, immediately on reduction of the temperature from  $565^{\circ}$  to  $228^{\circ}$ , the value of  $k$  was 0.0036. After cooling at room temperature for some time,  $k$  had fallen to 0.0026. The operation of both effects may be seen in Series 14, 15, and 16. Thus, the value of  $k$  at  $470^{\circ}$  on the first day after cooling from  $530^{\circ}$  was 0.0660, but had fallen by the second day to the normal value of 0.0560. On further cooling to  $373^{\circ}$ ,  $k$  followed an analogous course, having the values 0.0225 on the first, and 0.0200 on the second day after such reduction in temperature. When the temperature was again increased to  $485^{\circ}$ ,  $k$  had the abnormally low value of 0.0570 on the first day.

(5) The general facts relating to the electrical condition of the gauze established by the results recorded in Table I are as follows. (a) The potentials acquired in the moist system differed from those in the dry at the same temperature (see Series 8 followed by 9, and cf. also 1 and 3), (b) the difference between the potentials in the moist and dry tended to decrease as the temperature was increased.

(6) The coefficient of charging rate, PC, did not, in general, remain constant at constant temperature. (a) when the temperature of the surface had been increased from a lower value, PC diminished gradually at constant temperature. At low temperatures in either system, the fall was rapid at first (see Series 3

and 4), but PC soon attained a practically constant value. At high temperatures, PC fell steadily throughout the period of heating, a decrease parallel to the decrease in the value of  $k$  in the same conditions (see Series 6, 7 and 12). In Series 13 at  $530^\circ$ , the values of PC and  $k$  passed simultaneously through a maximum. At low temperatures, however, a rise in  $k$  was in certain cases accompanied by a fall in PC (see Series 1, 3 and 4), but these three may all be regarded as anomalous in some particular. In the only case where PC gradually rose to constancy (Series 5), it was accompanied by a parallel rise in the value of  $k$ . (b) On reduction of the temperature of the surface, PC at first rose rapidly (see Series 14 and 15). In Series 8, the value of PC was abnormally low immediately after cooling to  $228^\circ$  from a higher temperature, after the surface had been cooled to room temperature, however, and then re-heated to  $228^\circ$ , PC rose to the exceptionally high value, 0.7, but on continued heating this fell to the normal value of 0.2 for that temperature. (c) For a given temperature, the value of PC in the moist system was lower after the period of drying than before. In both moist and dry systems, the value of PC passed through a maximum at the temperature where the sign of  $A$  changed from positive to negative.

#### *Further Experimental Results relating to the Catalytic Reaction*

*Variation of the Reaction Velocity with the Gaseous Pressure*—Throughout the experiments, in either the moist or the dry system, it was found that during the combustion of a given charge of the gaseous mixture, the rate of reaction did not accurately follow a unimolecular law, the rate of decrease of pressure falling off less rapidly than required by such a law. In consequence, the curves connecting  $\log p$  and  $t$  were not linear except over small ranges of pressure. The extent of the variation is shown in fig. 3, where  $k$  (measured by the slope of the tangent to the  $\log p, t$  curve) is plotted against pressure.

Experiments were carried out on the rate of combination of  $2\text{CO} + \text{O}_2$  mixtures on the surface of a heated gold wire, in order to determine whether the form of the surface affected the type of variation of  $k$  with pressure, with negative results. It is therefore clear that the effect of pressure cannot be attributed to the operation of diffusion factors dependent on the use of a catalyst in the form of gauze.

*Effect of Evacuation on the Reaction Velocity*—In either the dry or the moist system, evacuation resulted in a reduction of the reaction velocity when the  $2\text{CO} + \text{O}_2$  mixture was subsequently admitted. Such reduction was, however, only temporary and the surface soon regained its normal state of activity.

corresponding to the conditions under investigation. The recovery was complete within three hours at 250° in the moist system, in 24 hours in the dry, and within a few minutes in either system at 450° or above.

*Catalytic Combustion on the Walls of the Quartz Vessel*—The rates of reaction in the dry and moist systems between 225° and 650° in the quartz vessel from which the gold surface had been removed are shown in Table III. An important feature of these results is that they show that the value of  $k$  was throughout higher in the dry than in the moist system.

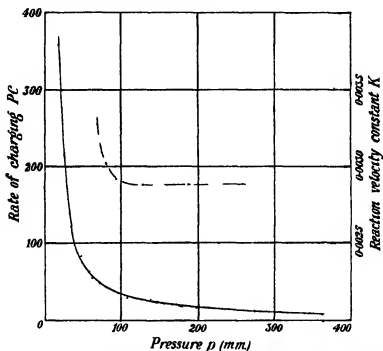


FIG. 3 — — —  $k \propto p$  (Series 2, 334° C), ———  $PC \propto p$  (Series 5, 495° C)

Table III

<i>Dry System</i>						
Temperature °C	226	333	457	571	602	—
$k$	0.0000	0.0000	0.0001	0.0047	0.0063	—
<i>Moist System</i>						
Temperature °C	225	350	502	550	587	580
$k$	0.0000	0.0000	0.0001	0.0002	0.0007	0.0012

*The Equilibrium Potential of the Surface*—For purposes of comparison, the electrical condition of the surface was examined in the individual gases CO,



$O_2$  and  $CO_2$ , both moist and dry, throughout the temperature range covered by the experiments on reaction velocity, the results being summarized in fig 4

The sign and magnitude of the equilibrium potential of the surface in the reacting mixture varied considerably in the moist system at temperatures below  $400^\circ$  (see Table I) No such variations were observed with any of the individual gases in the moist, or in the dry in any conditions. The magnitude of A was found to be independent of the gaseous pressure throughout the range covered in the experiments

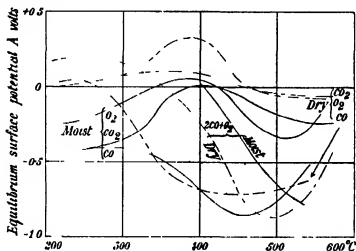


FIG 4

The following facts are evident from fig 4 —

- (1) All the curves showing the variation of A with temperature have a general formal similarity
- (2) In the dry system (a) The  $2CO + O_2$  curve approximated more closely to the CO curve than to that for  $O_2$ . At lower temperatures, the value of A for the mixture had a higher positive value than that for either CO or  $O_2$  or their sum, passed through zero at the temperature where the potentials in CO and  $O_2$  began to diverge considerably, became equal to the CO A-value at the temperature where the  $O_2$  A-value was zero, and rose to a higher positive value than CO in the range of temperature in which both CO and  $O_2$  had negative A-values (b) The  $CO_2$  curve followed the  $O_2$  curve closely throughout, the magnitude of A in  $CO_2$  being, however, uniformly less than in  $O_2$ .

- (3) In the moist system (a) The  $2\text{CO} + \text{O}_2$  curve followed the  $\text{O}_2$  curve in general form more nearly than that for  $\text{CO}$  (b) The  $\text{CO}_2$  curve approximated to the  $\text{CO}$  curve at low temperatures and to the  $\text{O}_2$  curve at high temperatures
- (4) On comparing the curves for the individual gases in the moist and dry systems respectively, it will be seen that (a) the curves for all three gases were displaced in a negative direction by the presence of moisture, (b) the  $\text{CO}$  and  $\text{CO}_2$  curves were displaced more at low than at high temperatures, and, finally, (c) the  $\text{O}_2$  curve was displaced uniformly throughout, and showed signs of a change of form at high temperatures in the moist
- (5) The values of  $\lambda$  for the reacting mixture  $2\text{CO} + \text{O}_2$  were displaced uniformly in a positive direction by the introduction of water

*The Rate of Electrical Charging of the Surface* - The rate of charging of the gold surface increased rapidly with decreasing pressure, the type of increase which occurred is shown in fig 5 All values of  $\text{PC}$  quoted in the present communication refer to a total gaseous pressure of 100 mm

The experiments on the rate of charging ( $\text{PC}$ ) in the reacting mixture, and in the individual gases are summarized in Table IV

Table IV

Gas		250° C	300° C	350° C	400° C	450° C	500° C	550° C
Moist	$\text{O}_2$	6.5	6.5	7.9	13	32	156	1750
	$\text{CO}$	11	19	33	59	130	500	8000
	$\text{CO}_2$	30	53	100	282	708	2000	6100
	$2\text{CO} + \text{O}_2$	1.6	1.6	2.0	3.5	7.6	45	216*
Dry	$\text{O}_2$	0.0	1.0	2.0	11.5	42	—	—
	$\text{CO}$	0.8	1.6	5.0	32	400	1510	2510
	$\text{CO}_2$	—	—	0.4	2.5	25	200	1280
	$2\text{CO} + \text{O}_2$	0.37	0.50	0.45	0.25	1.9	45	180

\* At 530°

The following facts are clearly brought out in Table IV —

- (1) Increase of temperature was not invariably accompanied by an increased rate of charging. Thus, the value of  $\text{PC}$  for the dry, reacting mixture  $2\text{CO} + \text{O}_2$  fell from 0.50 to 0.25 as the temperature was raised from 300° to 400°.  $\text{PC}$  remained constant in the moist mixture while the temperature was raised from 250° to 320°, and in moist  $\text{O}_2$  over the

temperature range from 200° to 300° In all three cases the temperatures were in the neighbourhood of that at which the sign of  $A$  changed in the respective systems

- (2) In general, the rate of charging in the moist was much greater than in the dry system, but the differences became less marked as the temperature was increased

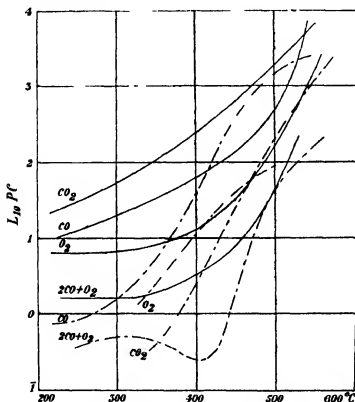


FIG 5 ——— Moist series, — — — dry series

- (3)  $CO$  was an exception to the general rule that the rate of charging was greater in the moist than in the dry. Over the range of temperature 420° to 530°,  $PC$  was higher in the dry than in the moist, although at higher and lower temperatures the reverse occurred. The  $PC$  values for the reacting mixture became identical at 480°, whilst diverging at higher and lower temperatures
- (4) The  $PC$  values for  $O_2$  in either system were considerably less than those for  $CO$  and  $CO_2$ , although greater than those for the reacting mixture  $2CO + O_2$

In order to exhibit the effect of moisture on the rate of charging more clearly the graphs of  $\log PC$  plotted against temperature are shown in fig 5. In addition to those set forth above, the results incorporated in fig 5 establish the following facts —

- (1) The slopes of the curves for the individual gases and the reacting mixture in the moist system increase steadily with increasing temperature, while those for the dry system pass through one or more points of inflexion.
- (2) The mean rate of increase of  $PC$  with temperature is throughout higher in the dry than in the moist system, an observation which may be compared with the higher rate of increase of reaction velocity,  $k$ , with temperature in the dry, compared with that in the moist system.

#### *Variation of Reaction Velocity with Temperature*

In fig 2 the values of  $\log k$  are shown plotted against the reciprocals of the corresponding absolute temperatures. The dry and second moist series give linear relationships over the greater part of the temperature range. In the former series, deviations occur at the higher temperatures where the gradual decrease of activity previously noted took place (Series 6 and 7). In the second moist series, the points representing Series 9 and 13 both lie above the line joining the remainder of the points. The former may represent a change in the heat of activation at low temperatures, but is more probably to be ascribed to the other changes in the surface brought about by the long period of drying. The latter deviation appears to represent a change in the course of the reaction at the highest temperatures attained. The results for the reaction on the surface of the quartz vessel (Table III) show that it cannot be due to a change from heterogeneous to homogeneous reaction.

The slopes of the graphs give values for the apparent heats of activation of 12,000 and 8730 cal per gm mol for the dry and second moist series respectively. The slope of the line joining the points corresponding to the two series of experiments in the first moist series gave a value of 8500 cal per gm mol. It may therefore be concluded that the period of drying did not change the mechanism of the moist reaction on the gold surface.

The apparent heat of activation of a heterogeneous reaction such as that under consideration is a composite quantity, representing not only a true heat of chemical activation, but also additional energy factors connected with the processes of adsorption and desorption on the surface. In the absence of sufficient data relating to the adsorption of  $CO$ ,  $O_2$  and  $CO_2$  by gold in moist

and dry systems, the differing heats of activation in the moist and dry cannot be taken as direct proof that the mechanisms of the two reactions are dissimilar. It cannot be supposed, however, that the presence of moisture would modify the processes of adsorption and desorption of gaseous molecules on the surface and leave the chemical reactions unaffected.

### Discussion

The results set forth above establish the facts that under the conditions of our experiments the heterogeneous catalytic combustion of carbonic oxide by oxygen in contact with a gold surface approximates more closely to a reaction of the first, than to one of any other order, and that the reaction is not retarded by the presence of carbon dioxide. Bone and Andrew\* have, however, previously shown that, under conditions similar to those employed by us, neither carbonic oxide nor oxygen are strongly adsorbed by the catalysing surface. The conclusion may, therefore, be drawn that the order of reaction found experimentally is also, in fact, the true order, and that the rate of combustion of a  $2\text{CO} + \text{O}_2$  mixture in contact with gold is directly proportional to the concentration of one of the two constituents of the mixture, probably carbon monoxide. The fact that  $k$  increased with falling pressure can then be reasonably ascribed to an increase in the value of the ratio of adsorbed to free gaseous carbon monoxide.

The remarkable deactivating effect of evacuation upon the catalytic powers of the gauze, also previously observed by Bone and Andrew (*loc. cit.*), appears to have its parallel in a recent observation by Finch and co-workers† to the effect that the catalytic activity of a sputtered platinum film may be completely destroyed by drying or by evacuation. It would seem that drying results in the growth of larger crystal aggregates with consequent closing up and elimination of Smekal cracks. The fact that the activity of the previously dried gold surface can be relatively rapidly restored by heating in contact with  $2\text{CO} + \text{O}_2$  strongly suggests that activation of, and interaction between these gases at the surface resulted in a reopening of fresh Smekal cracks.

The anomalous increase in the reaction velocities at the highest temperatures in the moist system occurred within the temperature range where the rate of the dry reaction closely approached that of the moist reaction. In view of the above considerations this fact suggests that at such temperatures both the dry and moist reaction mechanisms occurred simultaneously and independently,

\* 'Proc. Roy. Soc.,' A, vol. 106, p. 459 (1925)

† 'Proc. Roy. Soc.,' A, vol. 141, p. 414 (1933)

and, furthermore, that the nature of surface structure most favourable for promoting the dry reaction differed greatly from that suitable in the combustion of the moist mixture. Thus, on changing from the dry to the moist reaction or *vice versa*, a more or less prolonged period of induction, amounting, in one case, to 28 days, elapsed before the surface had become completely normalized to the changed conditions of reaction. These facts strongly suggest that the mechanisms of the moist and dry reactions differ, and it may be reasonably supposed that in the dry reaction adsorbed oxygen is burnt directly by carbonic oxide, whereas, in the presence of moisture the carbonic oxide reacts with adsorbed steam, the hydrogen thus liberated reacting with adsorbed oxygen.

The close connection between the structure and catalytic properties of a surface has been previously recognized by Bone, Langmuir, Rideal, Taylor and others, and the above experiments enable us to discriminate between two processes resulting in profoundly different classes of surface structural change. Of these, normalization towards a specified reaction or temperature is one type, the other being a process of sintering whereby the activity of the surface, as measured in terms of the rate of a standard reaction, was permanently unpaired.

Reaction and temperature normalization of the surface differed in themselves in that normalization at an elevated temperature left the surface in a temporarily abnormally active condition on return to a lower temperature. After sufficiently prolonged promotion of the reaction, however, the activity of the surface returned to the normal value. These facts in conjunction with those previously discussed suggest that heating, or the promotion of either the moist or the dry reaction all lead to the formation of different types of surface structure. Sintering, on the other hand, which must be regarded purely as a temperature effect, produces structural changes deleterious to catalytic action from which complete recovery was not possible under the conditions of our experiments.

The electrical properties of the gold gauze in different conditions show clearly that the influence of water cannot be regarded as merely additive, since, if such were so, its effect, as revealed by the values of  $A$  and  $PC$ , would be the same for the three individual gases as for the reacting mixture, whereas the experiments have shown the effect to be specific. This is particularly in evidence in the values of  $A$  for the reacting mixture, in which a positive displacement of potential occurs on introduction of water, contrasted with the negative displacement brought about in all the individual gases. It is clear that, during the progress of chemical reaction, some molecular complex or type of configuration was present on the surface of the metal which was not capable

of being formed by contiguous adsorption of water and either  $\text{CO}$ ,  $\text{O}_2$  or  $\text{CO}_2$ . While this fact alone does not prove that water participated in the reaction, the close relation which was found to exist between the rate of electrical charging and the rate of chemical reaction makes it difficult to resist such a conclusion.

Further support for the view that water plays an essential part in the surface reactions may be derived from the observations on the rate of charging. It has been shown that the introduction of water not only changed the general type of functional relationship between the rate of charging and temperature, but that the different gases were not affected to the same extent. In both moist and dry systems, the rate of charging during chemical reaction was considerably less than in any of the individual gases. Since, in general, the energy necessary to ionize a molecule is less than that required to excite it to a state of chemical activity, it may be supposed that the probability of ionization of molecules adsorbed on the gold surface was substantially reduced by the occurrence of chemical reaction. The product of such reaction ( $\text{CO}_2$ ) must, at least for a short time after its formation, have been in a favourable condition for ionization. Of all the individual gases,  $\text{CO}_2$  had the lowest rates of charging in the dry, and the highest in the moist. It would, therefore, be anticipated that the differences between the rate of charging in the reacting mixture and that in either  $\text{CO}$  or  $\text{O}_2$  would be less in the moist than in the dry, a conclusion which is borne out by the experiments.

In only one case did the rate of charging attain a higher value in the dry system than in the moist, namely, in  $\text{CO}$  over the temperature range  $420^\circ$  to  $530^\circ$ . This anomaly may be attributed to the occurrence of the water gas reaction in the moist system and a consequent reduction in the rate of charging. The marked resemblance previously pointed out between the electrical properties of the surface in  $\text{CO}$  and in the reacting mixture over this range of temperature lends further support to the view that oxidation of  $\text{CO}$  in the moist system takes place indirectly through the intervention of adsorbed water.

The experimental results set forth above have shown that, in general, when the surface was fully normalized at a constant temperature, gradual changes in the value of the reaction velocity constant,  $k$ , were accompanied by parallel changes in the coefficient of charging rate,  $PC$ . When, on the other hand, the surface of the metal was passing through readjustments resulting from changes in temperature,  $k$  and  $PC$  were no longer parallel. In particular, the rapid initial rise or fall in the value of  $k$  following increase or decrease of temperature was usually accompanied by change of  $PC$  in the contrary sense, decrease in

rate of charging resulting from increase of temperature and *vice versa*. Following the view of structural changes of the surface put forward above, it is clear that the extent to which activation or ionization of molecules adsorbed on the surface occurs must depend upon the simultaneous operation of two factors, definable as accessibility and suitability respectively. Sintering of the surface, which has been characterized in the foregoing as an irreversible decrease in catalytic activity, is to be regarded as reducing the accessibility of catalytic surface, and in the present experiments led to decrease of both the rate of chemical reaction and the rate of electrical charging. Factors, other than sintering, upon which the catalytic activity of the surface depends, may be classified as suitability factors, distinguishable by the period of time required for their operation which was short compared with the long and gradual process of sintering. Two such factors have been elucidated in the present experiments, namely, reaction and temperature normalization. Reaction normalization the occurrence of which is seen most clearly in the moist Series 13 undoubtedly affects rate of reaction and rate of electrical charging in the same way. The operation of temperature normalization, on the other hand, which is made evident in the behaviour of the surface immediately after increase or decrease of temperature, leads to opposite changes in  $\lambda$  and PC. It is therefore clear that while, in general, the rate at which chemical reaction occurs on the surface is parallel to the rate at which the characteristic electrical interface potential is established, the magnitudes of the two rates are determined to some extent by forces which act in contrary directions.

We are carrying out further investigations on the nature of these forces.

One of us (B W B) wishes to thank the Department of Scientific and Industrial Research for a grant and also the Trustees of the Beit Fellowship Fund for a Fellowship during part of the tenure of which this work was carried out.

### *Summary*

Previous experimental study of the electrical condition of heated metals having indicated the existence of a striking analogy between such electrical condition and the catalytic properties of the surfaces, a series of experiments with a gold gauze surface was carried out in such a manner that the catalytic and electrical activities of the metal could be simultaneously observed and followed. The reaction selected was the heterogeneous combination of carbon monoxide and oxygen, in view of the well-known influence of water on this reaction, series of observations were made in both moist and dry systems, with



the further object of elucidating the function of water in the reaction. The electrical condition of the metallic surface was expressed in terms of the magnitude and sign of the equilibrium potential which it acquired in given conditions, and its electrical activity was measured by the specific rate at which that potential was approached on insulation at zero or other standard potential.

In similar experimental conditions, the reaction velocities were invariably higher in the moist than in the dry reaction system, with increasing temperature, however, the differences between the rates observed in the two systems diminished practically throughout the range of temperature covered by the experiments. Several types of gradual change in the catalytic activity of the surface were observed, some reversible, others permanent. Such changes were ascribed to alterations in the superficial structure of the metal and were correlated with the effects of temperature, evacuation of the catalyst, and the presence or absence of moisture on the surface.

In general, throughout the experiments, changes in the rate of electrical charging of the metal followed closely the corresponding changes in the catalytic activity, increasing with rising temperature or with the introduction of water, and undergoing similar variations to the rate of reaction when the surface was maintained at constant temperature. The equilibrium potential of the metal gave further evidence of the close connection between the electrical and catalytic properties of the surface, being affected by the presence of moisture and by the occurrence of chemical reaction. The evidence derived from the electrical properties of the surface indicated that water played an essential part in the changes which occurred in the layer adsorbed on the metallic surface.

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### *The Crystal Structure of the Heusler Alloys*

By A J BRADLEY, D Sc., Royal Society Warren Research Fellow,  
and J W RODGERS, B Sc

(Communicated by W L Bragg, F R S -- Received October 4, 1933)

Beginning in the year 1898 Heusler\* discovered a series of ferromagnetic alloys the most important containing copper, manganese, and aluminium. They are characterized by remarkable magnetic properties, because although composed only of paramagnetic or diamagnetic elements, they become ferromagnetic after suitable heat treatment †. Various explanations of this property have been advanced, but it was usually considered to be due to the formation of a series of solid solutions of the type  $(\text{CuMn})_3\text{Al}$ , in which the proportions of copper and manganese may be varied within fairly wide limits.

The Heusler alloys have been repeatedly investigated by means of X-rays. Young, ‡ using molybdenum radiation examined alloys of two different compositions. He found that one was face-centred cubic, while the other was a mixture of face-centred and body-centred cubic structures. The mixed alloy was the more magnetic.

A more detailed investigation was made by Leiv Harang, § using copper radiation. He found three structures, face-centred cubic, body-centred cubic, and a structure similar to that of  $\gamma$  brass, which correspond respectively to the  $\alpha$ ,  $\beta$ , and  $\delta$  phases of the copper-aluminium system ||. They are successively produced by increasing the proportion of aluminium in the alloy. Harang did not find it possible to trace any relation between the magnetic properties and the crystal structure, and therefore could not ascribe them to a single lattice. These conclusions are not in agreement with recent investigations ¶.

Later, Elis Persson investigated these alloys by means of chromium radiation. In a preliminary note\*\* he showed that the structure of a ferromagnetic alloy

\* Heusler, Stark, and Haupt, 'Verh. deutsch. phys. Ges.', vol 5, p 219 (1903), Heusler and Richarz, 'Z. anorg. Chem.', vol 61, p 269 (1908).

† Take, 'Inaug. diss. Marburg' (1904).

‡ 'Phil. Mag.', vol 46, p 291 (1923).

§ 'Z. Kristallog.', vol 65, p 261 (1927).

|| Jette, Westgren, and Phragmén, 'J. Inst. Metals', vol 31, p 201 (1924), Obinata, 'Mem. Ryojun Coll. Eng.', vol 31, pp 3, 286, 295 (1929), Bradley and Jones, 'J. Inst. Metals', vol 51, p 131 (1933).

¶ Krings and Ostmann, 'Z. anorg. Chem.', vol 163, p 154 (1927), Heusler, 'Z. anorg. Chem.', vol 171, p 126 (1928).

\*\* 'Naturwiss.', vol 16, p 613 (1928).

corresponding to the formula  $\text{Cu}_2\text{MnAl}$  was body-centred cubic, with the aluminium atoms forming a face-centred superlattice. The structure was therefore like that of  $\text{Fe}_3\text{Si}^*$ , or that of  $\text{Fe}_3\text{Al}^\dagger$  discovered later. The unit cell is built up of eight small body-centred cubes, and therefore contains 16 atoms, of which four are aluminium, four manganese, and the remainder copper. The aluminium atoms form a face-centred cube with double the dimensions of the small body-centred cube. No attempt was made to find the position of the manganese atoms by means of X-rays.

Potter<sup>‡</sup> made an X-ray examination of single crystals using copper radiation. He believed that the Mn atoms occupied special positions like the Al atoms. On account of the resemblance between the magnetic properties of this alloy and nickel, he concluded that its ferromagnetism was due to the manganese atoms being arranged on a face-centred cubic lattice.

In continuation of his earlier researches, Persson<sup>§</sup> came to the same conclusion. The series of alloys  $(\text{CuMn})_2\text{Al}$  is only ferromagnetic when the manganese content exceeds 19% (atomic). This is independent of the arrangement of aluminium atoms, which is apparently the same whatever the amount of manganese present, within wide limits. Persson believes that the presence of 19% of manganese is required in order to produce a regular arrangement of manganese atoms.

Persson attempted to find whether the special  $\text{Cu}_2\text{MnAl}$  type of structure was alone responsible for the ferromagnetic properties of the copper-manganese-aluminium alloys. He examined a whole series of alloys of the composition  $(\text{CuMn})_2\text{Al}$ , varying the proportions of copper and manganese. An X-ray examination was made after heat treatment of the powdered alloy. In one series of experiments the alloys were quenched in water from a temperature only  $50^\circ$  below the melting point. In another series the alloys were tempered for 350 hours at  $210^\circ$ .

The quenching experiments showed that four different types of structure occurred. Face-centred cubic, body-centred cubic, and 'Gamma' structures corresponded to the  $\alpha$ ,  $\beta$  and  $\delta$  phases of the copper-aluminium system, and a fourth phase possessed a structure like that of  $\beta$  Mn. Contrary to Haraug, Persson concluded that only the body-centred cubic  $\beta$  phase was ferromagnetic, the ferromagnetism of the alloys increasing with the amount of the  $\beta$  phase present.

\* Phragmén, 'Tekn. Tidskrift' (Stockholm), vol. 56, p. 81 (1926).

† Bradley and Jay, 'Proc. Roy. Soc., A', vol. 136, p. 210 (1932).

‡ 'Proc. Phys. Soc.', London, vol. 41, p. 135 (1929).

§ 'Z. Physik', vol. 57, p. 115 (1929).

The tempering experiments showed that the  $\beta$  phase was decomposed by this form of heat treatment. The alloy splits up into two or more phases. If the Mn content is less than required for the formation of the alloy  $\text{Cu}_2\text{MnAl}$ , the phases are  $\text{Cu}_2\text{Al}$  with lattice spacing 5.833 Å, and  $\text{Cu}_2\text{MnAl}$  with lattice spacing 5.950 Å. If the Mn content is greater, a structure of the  $\beta$  Mn type with lattice spacing 6.370 Å is formed, together with  $\text{Cu}_2\text{MnAl}$  with lattice spacing 5.950 Å. Maximum ferromagnetism corresponds to the greatest amount of  $\text{Cu}_2\text{MnAl}$ .

Whether the alloy is quenched or tempered the lattice spacing of the  $\beta$  body-centred cubic phase never exceeds 5.950 Å. For quenched alloys the lattice spacing increases linearly with increase of manganese content between  $\text{Cu}_2\text{Al}$  and  $\text{Cu}_2\text{MnAl}$  and ferromagnetism is the greater the greater the lattice spacing.

An alloy of the composition  $\text{CuMnAl}_2$  has the CsCl type of structure and is non-magnetic. The property of ferromagnetism is thus associated in a peculiar degree with the special composition  $\text{Cu}_2\text{MnAl}$ . The closer the alloy attains to this composition the more marked is its ferromagnetic character. Pearson concludes that the property of ferromagnetism is due to the nature of the crystal structure of  $\text{Cu}_2\text{MnAl}$ , and is linked up with the mode of distribution of the manganese atoms.

The object of the present paper is to fix the position of the manganese atoms in the magnetic alloys by direct experiment, and to test whether a change of structure *without change of composition* will destroy the ferromagnetic character of the alloy. This should decide whether structure or composition is the more important condition for ferromagnetism.

### I Present Experiments

Eight alloys of the approximate composition  $\text{Cu}_2\text{MnAl}$  were prepared by melting together 50 gms. of copper, manganese, and aluminium in slightly different proportions in an alumina lined crucible\* in a high frequency induction furnace, under a low pressure of hydrogen. In order to remove coring, and to make the alloys homogeneous, they were heated for 6 hours at 750° C in an electric furnace and allowed to cool slowly down to room temperature. Drillings were then taken by means of a special drill,† kindly supplied by Mr Gardiner, of Easterbrook Allcard & Co., Sheffield. The drillings gave small particles which could be ground in an agate mortar. The fine powders so obtained were

\* Jay, 'J. Iron and Steel Inst.', vol. 125, p. 427 (1932).

† Edgar Allen's Stag Major.

sieved through a mesh 250 to the inch, and then subjected to further heat treatment

A portion was heated in hydrogen at 500° C for 6 hours, and then allowed to cool slowly down to 300° C over a period of a few hours. At this point the current was switched off, and the powders were allowed to cool to room temperature in the furnace. X-ray powder photographs of these alloys were taken using radiation from an iron anticathode. They show that the bulk of each alloy consists of a phase of the  $\gamma$  brass type\* like  $\text{Cu}_3\text{Al}_4$ † and that at least one other phase is present.

Very different photographs were obtained from the second portions of the powders. These were heated in hydrogen to a temperature of 800° C and after about half an hour quenched by allowing a stream of cold water under pressure to enter the furnace. The structure is now body-centred cubic with a superlattice of the typical Heusler alloy type, as described by Persson and Potter. These photographs were repeated using radiation from copper and from zinc anticathodes. There are differences between the relative intensities of the fainter lines of the three photographs, which we shall explain later.

In the alloy selected for detailed investigation the proportions of the ingredients did not correspond exactly to the theoretical values for the composition  $\text{Cu}_2\text{MnAl}$ . Analysis‡ of the alloy showed that the approximate composition was Cu 67.5%, Mn 17.5%, Al 15% corresponding to the atomic composition Cu 2.2, Mn 0.65, Al 1.15. The deficit of manganese was therefore made up partly by copper and partly by aluminium. This particular alloy was chosen because it was the only one which showed an almost complete change of structure with the two methods of heat treatment. After annealing at 500° and slowly cooling down to room temperature, it gave an X-ray powder photograph corresponding to the  $\delta$  copper aluminium structure ( $\text{Cu}_3\text{Al}_4$ ), with only a faint trace of lines belonging to another pattern. In this state the alloy was found to be practically non-magnetic. On the other hand, the same powder after quenching from 800° showed only a body-centred cubic structure with face-centred superlattice. The alloy was now strongly ferromagnetic.

\* Westgren and Phragmén, 'Phil Mag.', vol 50, p 331 (1925), Bradley and Thewlis, 'Proc Roy Soc.' A, vol 112, p 678 (1926).

† Bradley, 'Phil Mag.', vol 6, p 878 (1928), Bradley and Jones, 'J Inst Metals,' vol 51, p 131 (1933).

‡ The alloys were analysed by Mr J. W. Cathbertson of the University, to whom the authors are indebted for his kindness.

The above facts show conclusively that the magnetic properties of the alloy are dependent on the type of crystal structure, as Persson suggested. The composition of the alloy after heat treatment was checked by chemical analysis, and found to be unaltered\*. There is therefore no doubt that the ferromagnetism of the alloy is not a matter of chemical composition but of atomic arrangement.

The arrangement of the atoms in the annealed and slowly cooled alloy can be fixed to some extent from a visual inspection of the photograph. Its resemblance to the powder photograph of  $\text{Cu}_3\text{Al}_4$  is extraordinarily close. In  $\text{Cu}_3\text{Al}_4$ , the unit cell is cubic and contains 52 atoms of which 36 are copper

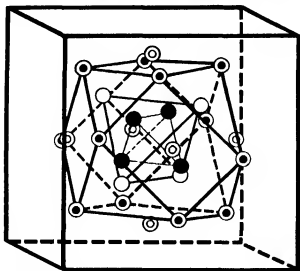


FIG. 1—Annealed Heusler alloy (non magnetic), ● A, ○ B, ⊙ C, ⊗ D

and 16 aluminium. The copper and aluminium atoms each occupy definite positions in the lattice. From considerations of symmetry the 52 atoms can be divided into eight sets which may be called  $A_1, A_2, B_1, B_2, C_1, C_2, D_1, D_2$  respectively. Four  $A_1$ , 4  $B_1$ , 6  $C_1$ , and 12  $D_1$  constitute a cluster of 26 atoms which are grouped symmetrically around the centre of the unit cell, fig. 1. A second cluster of 26 atoms, not shown in the figure, is grouped symmetrically about each corner of the cell, so that the whole structure would be body-centred cubic, except for the fact that there are more copper atoms and fewer aluminium atoms in one cluster than in the other. The aluminium atoms are concentrated in  $A_1$  and  $D_2$ . The copper atoms occupy the remaining positions.

\* The manganese contents of the quenched and annealed specimens agreed to 0.1%.

The similarity of the powder photographs of the annealed Heusler alloy and that of  $\text{Cu}_2\text{Al}_4$  shows at once that the atomic co-ordinates are almost identical in the two. Further, it can be concluded that the aluminium atoms again occupy positions  $A_1$  and  $D_2$ . The remaining positions are filled chiefly by copper atoms, but about one-fourth of these are replaced by manganese. The formula of the annealed alloy can therefore be written as  $(\text{CuMn})_8\text{Al}_4$ . The proportions of aluminium are not quite sufficient to satisfy this formula, which may possibly explain the slight admixture of the second phase, shown by the powder photograph. Without a more detailed investigation, it is impossible to say whether the manganese atoms occupy special positions in the lattice, or whether they are mixed up with the copper atoms in a purely random manner, but there is no doubt about the aluminium atoms.

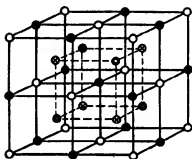


FIG 2—General type of structure,  
○ A, ⊙ B, ● C, ⊗ D

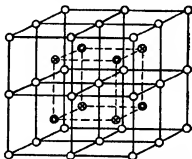


FIG 3—Quenched Heusler alloy (magnetic), ○ Cu, ⊙ Al, ⊗ Mn

The ferromagnetic quenched alloy has a unit cell containing 16 atoms, fig 2. These consist of four sets of atoms (A, B, C, D) each corresponding to a face-centred cubic lattice. The superlattice lines are caused by the segregation of the aluminium atoms into one of the four sets of positions (say B). Both Persson and Potter suggest that the ferromagnetism of these alloys is due to the manganese atoms occupying a special position as in fig 3. It was our object to test this hypothesis. The question is whether the manganese atoms are mixed up at random with the copper atoms or whether they keep to their own positions. The difference between the scattering powers of copper and manganese for X-rays is so slight that this might at first sight appear to be a matter of some difficulty, but, in fact, the problem has been solved by means of accurate photometer measurements of powder photographs taken with X-rays of different wave-lengths, making use of the anomalies in atomic scattering

factor which occur when the frequency of the radiation is close to the characteristic absorption frequency of the element

The powder photographs of the quenched alloys taken with iron, copper, and zinc radiations were photometered by means of a Cambridge micro-photometer. This instrument employs a null method, the blackening of the film being balanced against an Ilford wedge. Each film was calibrated by means of a rotating sector wheel, giving 10 uniform increments of intensity. On comparing the blackening of the film corresponding to each of the 10 steps, it is possible to trace the relationship between intensity and blackness characteristic of the film. For the films used in the present series of experiments the blackening curve is practically linear, so that the wedge readings may be taken as true measures of the intensity, some slight correction being applied to the strongest peaks where the calibration curve shows a slight departure from the linear law.

Table I—Observed Intensities Arbitrary Units

Line		Iron $K_{\alpha}$ radiation $\lambda = 1.934$		Copper $K_{\alpha}$ radiation $\lambda = 1.539$		Zinc $K_{\alpha}$ radiation $\lambda = 1.434$	
$\Delta h'$	$\Delta h$	Observed values	Corrected for absorption	Observed values	Corrected for absorption	Observed values	Corrected for absorption
3	111	110	96	119	171	123	158
4	200	330	236	172	198	152	161
8	220	4730	2040	4041	2750	3683	2490
11	311	78	26	92	48.5	67	36
12	222	171	53	71	34.5	49	24
16	400	1181	288	895	342	758	303
19	331	31	6.5	47	15.5	29	10
20	420	272	54	111	35	53	18
24	422	3974	694	2111	503	1616	462
27	511	58	8.5	35	8.5	43	11
32	333						
32	440						
35	531	2811	349	697	142	497	111
36	600	106	12	47	9	—	—
36	442	867	94.5	62	11	—	—

For each line wedge readings were taken at intervals of 0.1 mm., and the blackening values so found plotted against the distance along the film. The areas included under the peak curve and above the general level of the background intensity were computed, and tabulated in Table I. Corrections for the effect of absorption in the powder specimen were made by the method



of graphical integration described by Claassen \*. The values of the absorption factor depend on the product of the absorption coefficient ( $\mu$ ) and the radius of the specimen ( $r$ )

As a convenient method of support during the exposure, the specimen is diluted with Canada balsam and mounted on a hair. The effective value of  $\mu r$  is thereby reduced. The appropriate value of  $\mu r$  is most easily obtained from the following equation —

$$\mu r = \frac{\mu}{\rho} m \frac{1}{\pi r l}$$

The weight of the specimen ( $m$ ) was 0.0024 gm., the length ( $l$ ) 0.49 cm., and radius ( $r$ ) 0.0265 cm. The values of  $\mu/\rho$  were calculated with the help of Jonsson's tables,† for each of the elements Al, Cu, and Mn with each of the radiations Fe, Cu, and Zn. The results are summarized in Table II.

Table II

Radiation	$\lambda$	$\mu/\rho$ values				$\mu r$
		Al	Cu	Mn	Cu <sub>2</sub> MnAl	Cu <sub>2</sub> MnAl
Fe K <sub><math>\alpha</math></sub>	1.934	93.9	100	64	99.5	5.3
Cu K <sub><math>\alpha</math></sub>	1.539	51.2	50.4	285	113	6.7
Zn K <sub><math>\alpha</math></sub>	1.434	41.8	42.6	234	93.6	5.5

Applying these values of  $\mu r$  to the data given by Claassen, the values of the absorption factor were found for glancing angles  $0^\circ$ ,  $22\frac{1}{2}^\circ$ ,  $45^\circ$ ,  $67\frac{1}{2}^\circ$ , and  $90^\circ$ . Curves were then drawn giving the absorption factors for all angles between  $0^\circ$  and  $90^\circ$ , for each of the three radiations. From these curves were obtained the absorption factors by which the observed intensity values were divided.

The corrected intensity values in Table I are proportional to theoretical intensity values given by the equation

$$I = \frac{1 + \cos^2 2\theta}{\sin^2 \theta \cos \theta} p F^2,$$

where  $\theta$  is the glancing angle,  $p$  the number of co-operating planes, and  $F$  the structure factor at room temperature. With the help of this equation the

\* 'Phil. Mag.', vol. 9, p. 57 (1930).

† Jonsson, 'Uppsala Univ. Årskr.' (1928), Siegbahn, "Spektroskopie der Röntgenstrahlen," Julius Springer, Berlin (1931).

relative  $F$  values in Table III were obtained. They were reduced to a comparable scale by putting  $F_{220} = 100$  for each radiation.

This table shows some interesting features. On account of their smaller intensity, the superlattice lines  $\Sigma h^2 = 3, 4, 11, 12, 19, 20, 27, 35, 36$  are easily distinguished from the body-centred lines  $8, 16, 24, 32$ . The superlattice lines themselves may be divided into two series, the odd reflections  $3, 11, 19, 27, 35$  being definitely weaker than the corresponding even reflections  $4, 12, 20, 36$ . This distinction is to some extent obscured in Table I owing to the complications introduced by the  $\theta$  factor and the planar factor. It is shown better

Table III—Observed  $F$  Values for Three Radiations

Line		Iron $K_{\alpha}$ radiation $\lambda = 1.934$		Copper $K_{\alpha}$ radiation $\lambda = 1.539$		Zinc $K_{\alpha}$ radiation $\lambda = 1.434$	
$\Sigma h^2$	$hkl$	Relative $F$ values	Differences in neighbouring superlattice lines	Relative $F$ values	Differences in neighbouring superlattice lines	Relative $F$ values	Differences in neighbouring superlattice lines
3	111	15	} -17 {	17	} -8 {	18	} -6 {
4	200	12		25		24	
8	220	100		100		100	
11	311	10	} -15 {	11	} -6 {	10	} -5 {
12	222	25		17		15	
16	400	79		75		75	
19	331	6	} -12 {	9	} -5 {	8	} -2 {
20	420	18		14		10	
24	422	65		60		59	
27	{ 511 333 }	6	} — {	7	} — {	8	} — {
32	440	50		47		46	
35	531	4		6		—	
36	{ 600 442 }	11	} -7 {	8	} -2 {	—	} — {

by the figures given in Table III for the differences between the  $F$  values for neighbouring odd and even lines. The table shows that *the distinction between odd and even superlattice lines is far more marked with iron radiation than with copper or zinc radiation*. We shall now discuss the explanation of this phenomenon.

It will be shown that the differences with different radiations are due to the relation of the atomic scattering factor ( $f$ ) to the wave-length  $\lambda$ .  $f$  is usually given as a function of  $\sin \theta/\lambda$  which is independent of the wave-length, but this is not strictly correct.

## II The Atomic Scattering Factors of Aluminum, Manganese, and Copper

It has been shown in a number of experimental investigations\* that the atomic scattering factor ( $f$ ) of an element for X-rays depends upon the wave-length of the radiation, or to put the matter more precisely, the atomic scattering factor of an element is depressed by the use of radiation whose frequency lies close to the critical K absorption frequency of the scattering element. This fact was to be expected on theoretical grounds, and is analogous to anomalous dispersion in the optical region. It has been explained with the help of a combination of classical theory and quantum mechanics by Kallmann and Mark, Kronig, and Prins†. According to Coster and Knol,‡ the theory of Prins may be expressed in the following way:

The atomic scattering factor  $f$  of an element may be regarded as consisting of two components  $f_K$  due to the K electrons, and  $f_R$  due to the L, M, etc., electrons. Now in the present experiments, the factor  $f_R$  may as a first approximation be considered to be independent of the wave-length, because we are using radiation much harder than the critical L, M, etc., absorption edges.

On the other hand, for copper and manganese, the value of  $f_K$  is very susceptible to changes in wave-length, since all three radiations used are close to the K absorption edges of these elements.

According to Coster and Knol, for radiation on the short wave-length side of the absorption edge, there is a phase-change in the contribution of the K electrons to the atomic scattering factor. This may be expressed by writing  $f_K$  as a complex quantity, thus

$$f_K = f'_K + if''_K \quad (2)$$

where  $i = \sqrt{-1}$ , and  $f'_K$  and  $f''_K$  are two components with a phase difference of  $\pi/2$ , the resultant of which is  $f_K$ . The component  $f'_K$  either has the same phase as  $f_R$  or differs from it by an amount  $\pi$ . The component  $f''_K$  has a phase difference  $\pi/2$  compared with  $f'_K$  and  $f_R$ .

\* Mark and Saillard, 'Z. Physik,' vol. 33, p. 688 (1925), Armstrong, 'Phys. Rev.,' vol. 34, p. 931 (1929), Wyckoff, 'Phys. Rev.,' vol. 35, pp. 215, 583, 1116 (1930), Morton, 'Phys. Rev.,' vol. 38, p. 41 (1931), Coster, Knol and Prins, 'Z. Physik,' vol. 63, p. 345 (1930), vol. 75, p. 340 (1932), Glocker and Schäfer, 'Z. Physik,' vol. 73, p. 289 (1931), Bradley and Hope, 'Proc. Roy. Soc.,' A, vol. 136, p. 272 (1932).

† Kallmann and Mark, 'Ann. Physik,' vol. 82, p. 585 (1927), Kronig and Kramers, 'Z. Physik,' vol. 48, p. 174 (1928), Prins, 'Z. Physik,' vol. 47, p. 479 (1928), Kronig, 'Phys. Z.,' vol. 30, p. 521 (1929).

‡ 'Z. Physik,' vol. 75, p. 340 (1932), 'Proc. Roy. Soc.,' A, vol. 139, p. 459 (1933).

$f'_K$  is given by the equation

$$f'_K = n_K \left\{ 1 + \frac{\log_e |x^2 - 1|}{x^2} \right\}, \quad (3)$$

where  $x = \frac{\lambda_K}{\lambda}$ ,  $\lambda_K$  being the critical absorption wave-length of the scatterer, and  $\lambda$  the wave-length of the radiation

$f''_K$  is given by the equation

$$f''_K = \frac{\pi}{x^2} n_K \quad (4)$$

In each of the above equations  $n_K$  represents the full effect of the K oscillators away from the region of anomalous scattering. According to Prins the most probable value of  $n_K$  is 1.3, though Kronig and Kramers gave 0.86.

For radiation on the long wave-length side of the absorption edge there is no change of phase angle, and the expression for  $f_K$  therefore simplifies to

$$f_K = n_K \left\{ 1 + \frac{\log_e |1 - x^2|}{x^2} \right\}, \quad (5)$$

where  $n_K$  and  $x$  have the same meaning as before. Taken as it stands, equation (5), like equation (3), gives values of  $f = -\infty$  at the absorption edge, but Glocker and Schäfer have shown how this may be modified by the introduction of damping terms.

The change of phase angle on the short wave-length side of the absorption edge was verified qualitatively by the experiments of Coster, Knol, and Prins with zinc blende, and the general nature of the depression of the  $f$  values near the absorption edge is well verified by all the experimental work done in this region, but no satisfactory quantitative check has yet been obtained. According to most experimenters, the scattering factor is depressed on the short wave-length side considerably more than is demanded by theory. On the contrary, Bradley and Hope found that the scattering power of iron for copper  $K_\alpha$  radiation was very little depressed. Coster and Knol have recently pointed out that a correction must be made to this result to allow for the effect of a change in phase angle. However, this correction does little to bridge the difference from the results of other investigators, as may be seen from the following table.

We have chosen what we consider to be the most probable values of the scattering factors of copper and manganese, based on the experimental work of Bradley and Hope. They compared the values of the atomic scattering factor of iron for different radiations at the same value of  $\sin \theta/\lambda$ . For both

Table IV—The Atomic Scattering Factor of Iron ( $f_{110}$ )

Radiation	$\lambda/\lambda_K$	Glocker and Schäfer*		Wyckoff †	Bradley and Hope	Calculated from equations (3), (4), (5)
Mo $K_\alpha$	0.41	16.3	(17.1)	16.3	17.2	17.55
Cu $K_\alpha$	0.89	9.4	(11.6)	11.8	16.1 (corrected)	16.3
Ni $K_\alpha$	0.95	8.0	—	10.1	—	—
Co $K_\alpha$	1.01	—	—	—	14.1	12.6
Fe $K_\alpha$	1.10	11.9	(13.5)	13.8	15.7	14.5
Cr $K_\alpha$	1.31	12.7	—	—	17.2	15.25

\* Glocker and Schäfer have recently amended these values, and in a private communication to one of the authors they give values more nearly in agreement with those of Wyckoff. These are given in brackets.

† Corrected for temperature factor.

cobalt and iron radiations the scattering factor was found to be depressed below that for molybdenum radiation by an amount which was almost independent of  $\sin \theta/\lambda$ . This is to be expected on theoretical grounds. The depressions are due to changes in the contributions of the K electrons to the atomic scattering factor for different values of  $\lambda/\lambda_K$ . These depend only on  $n_K$  which is almost independent of  $\sin \theta/\lambda$ .

For copper  $K_\alpha$  radiation, according to Coster and Knol, the difference in scattering factor should be greater as the value of  $\sin \theta/\lambda$  increases, owing to the fact that  $f_K$  is a complex quantity. However, we may split up the atomic scattering factor into its two components, a real component ( $f_R + f'_K$ ) and an imaginary component,  $f''_K$  which differs in phase by  $\pi/2$ . At all values of  $\sin \theta/\lambda$ , ( $f_R + f'_K$ ) should be depressed by the same amount. It is therefore possible to obtain for each radiation a characteristic value of the depression of the "real" component of the  $f$  curve of iron, which is independent of  $\sin \theta/\lambda$ .

The depression in  $f$  in the neighbourhood of the K absorption edge is due to a variation in the contribution of the K electrons to the total  $f$  value of the atom. Whether we use a combination of classical theory and quantum theory (as Prins has) or a purely wave-mechanical theory,\* the variation in the contribution of the K electrons to the total value for the atom is a function of  $\lambda/\lambda_K$ .

It follows from this result that the anomalous dispersion by different elements can be deduced from experiments on one element only, provided the results

\* We are indebted to Dr. Williams of this University for discussing this matter with us.

are expressed in terms of  $\lambda/\lambda_K$ . Experiments are now being undertaken to test the validity of this generalization, the results of which will be communicated in due course.

In fig 4 the depressions in the observed  $f$  values for iron are plotted against the values of  $\lambda/\lambda_K$ ,  $\lambda$  being the wave-length of the radiations used (Mo, Cu, Co, Fe, Cr characteristic  $K_\alpha$  radiations), and  $\lambda_K$  the wave-length of the K

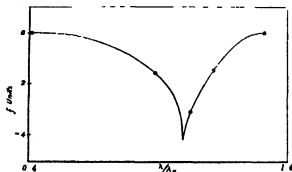


FIG 4.—Depression of  $f$  curves in the region of the K absorption edge.  $\circ$  Points from experiments on FeAl (Bradley and Hope) giving the difference between the normal  $f$  values and the values found near the absorption edge. For Cu radiation the value plotted is not  $f$  but  $f_K + f'_K$ .  $f''_K$  is calculated separately from the formula

$$f''_K = \frac{\pi}{\lambda^2} n_K$$

absorption edge of iron. The values for molybdenum radiation are taken as zero. In this figure the values of the depression at some distance from the absorption edge approach zero, and the biggest values of the depression are in the immediate neighbourhood of the K edge. Plotted as a function of  $\lambda/\lambda_K$ , theory indicates that the values represented on this curve should be universal. They represent the amounts to be subtracted from the normal  $f$  values where the frequency of the radiation is comparable with the critical absorption frequency.

In order to obtain the value of the atomic scattering factor for a given element with a given radiation for a given value of  $\sin \theta/\lambda$ , we first take the "normal" value of the atomic scattering factor of the element for the required value of  $\sin \theta/\lambda$ , and then subtract an amount read off from this curve, fig 4, to allow for the anomalous effects of the K absorption edge on  $f_K$ . Then we calculate the amount of  $f''_K$  from equation (4), assuming  $n_K = 1.3$ , and introduce a separate term into the calculations to allow for the effect of  $f''_K$  when

$\lambda/\lambda_K < 1$  An illustration will show how simple the procedure can be. To find  $f_{Mn} - f_{Al}$  for copper  $K_\alpha$  radiation we have

$$F = \sqrt{(f_{Mn} \text{ normal} - f_{Mn} \text{ depression} - f_{Al})^2 + f''^2_K} \quad (6)$$

Table V contains the necessary data for the evaluation of this expression, for the Heusler alloy

Table V —Depression in  $f$  Curve, due to Anomalous Dispersion

Scattering elements	$\lambda_K$	Zinc $K_\alpha$ radiation $\lambda = 1.434$	Copper $K_\alpha$ radiation $\lambda = 1.539$	Iron $K_\alpha$ radiation $\lambda = 1.934$
Al 13	7.936	0	0	0
Mn 25	1.892	1.0	1.0	3
Cu 29	1.377	2.5	1.5	0

Substituting in equation (6),

$$F = \sqrt{(f_{Mn} \text{ normal} - f_{Al} - 1.0)^2 + 2.7^2}$$

The normal  $f$  curves of aluminium, manganese, and copper were obtained from a paper by James and Brindley\*. The aluminium values were calculated by Hartree's method of self-consistent fields, for copper an approximate curve was obtained by the same method, for manganese, the Thomas values were used. It was shown by Bradley and Hope that the Thomas curve for iron gave an excellent agreement for molybdenum radiation, and since manganese is the next element to iron and does not differ greatly in atomic volume, it seems fairly safe to use the Thomas curve here, though, of course, it is probably less accurate than the other two calculated by Hartree's method.

It is now possible to understand the differences in the powder photographs taken with zinc, copper, and iron radiations. As Table V shows, the atomic scattering factor of manganese with iron radiation is three units less than normal, whereas that of copper with iron radiation is practically normal. Hence the difference in scattering power between copper and manganese is *three units greater than normal*. On the other hand, with zinc radiation the scattering power of copper is depressed more than that of manganese, so that the difference in scattering power between copper and manganese is now *1.5 units less than normal*. Thus iron radiation emphasizes the difference in scattering power between copper and manganese, whereas zinc radiation

\* 'Phil Mag.', vol 12, p 81 (1931), 'Z. Kristallog.', vol 78, p 470 (1931).

minimizes the difference. This fact naturally leads to differences in the intensities of the weaker reflections, where copper and manganese work in opposite directions. As we shall see later, the figures given in Table V provide not merely a qualitative explanation, but ultimately give a perfect quantitative explanation of all the observed intensities (Table VII).

On the other hand, if we had used the data of Glocker and Schafer or of Wyckoff, we should have been led to the conclusion that the scattering powers of copper and manganese with iron radiation ought not to be appreciably different from those with copper or zinc radiation. The different results obtained with different radiations prove that this is impossible. The theoretical expressions (equations (3), (4), and (5)) predict a difference between the scattering powers of copper and manganese for the different radiations employed, but do not explain the observed numerical values as well as our empirical values from the iron-aluminium experiments. Possibly more accurate results would be obtained if this theory took account of the variation in the contribution of the L electrons as well as the K electrons. The empirical curve, given in fig. 4, automatically makes this allowance, and partly for this reason should give accurate  $f$  values, if the experiments of Bradley and Hope are a sufficient basis for the curve. It is expected that new data will shortly become available, giving more points on the curve.

### III Possible Structure for the Heusler Alloy

The unit cell contains 16 atoms with the co-ordinates —

A	0 0 0,	$0 \frac{1}{2} \frac{1}{2}$ ,	$\frac{1}{2} 0 \frac{1}{2}$ ,	$\frac{1}{2} \frac{1}{2} 0$
B	$\frac{1}{2} \frac{1}{2} \frac{1}{2}$ ,	$\frac{1}{2} \frac{3}{4} \frac{3}{4}$ ,	$\frac{3}{4} \frac{1}{2} \frac{3}{4}$ ,	$\frac{3}{4} \frac{3}{4} \frac{1}{2}$
C	$\frac{1}{2} \frac{1}{2} \frac{3}{4}$ ,	$\frac{1}{2} 0 0$ ,	$0 \frac{1}{2} 0$ ,	$0 0 \frac{1}{2}$
D	$\frac{3}{4} \frac{3}{4} \frac{3}{4}$ ,	$\frac{3}{4} \frac{1}{2} \frac{1}{2}$ ,	$\frac{1}{2} \frac{3}{4} \frac{1}{2}$ ,	$\frac{1}{2} \frac{1}{2} \frac{3}{4}$

Symmetry considerations divide the atoms into the four sets A, B, C, D, as shown in fig. 2. The problem is to find which of these positions are occupied by copper, aluminium, and manganese atoms respectively. The data of Table III point the way to a solution.

Three facts must be explained —

- (1) The existence of three series of reflections, one strong series of lines [(1) 220, 400, 422, 440], and two weak series of superlattice lines

$$\left[ (2) 200, 222, 420, 600 \right], \left[ (3) 111, 311, 331, 511, 531 \right]$$



- (2) The superlattice lines of series (3) are definitely weaker than those of series (2)
- (3) The difference in intensity of the two series of superlattice lines depends on the wave-length of the radiation, being greatest for iron radiation and least for zinc radiation

Let the scattering powers in the four groups be  $f_a$ ,  $f_b$ ,  $f_c$ , and  $f_d$  respectively. Then the structure factors for planes of types (1), (2), (3) may be written in the following way —

$$(1) \quad f = f_a + f_b + f_c + f_d$$

$$(2) \quad f = (f_a + f_d) - (f_b + f_c)$$

$$(3) \quad f = \sqrt{(f_c - f_d)^2 + (f_a - f_b)^2}$$

From these equations it may be seen that the presence of reflections of both types (2) and (3) requires one group to be appreciably different in scattering power from the other three. Since copper and manganese are not very different in scattering power, this condition is equivalent to a requirement that most if not all the aluminium atoms should be sorted out into one group, say B.

If the atoms in groups A, C, D were equivalent in scattering power, reflections of types (2) and (3) would be equally strong. This is not so. The greater intensity of type (2) reflections shows that the scattering power of D groups is intermediate between those of A and C on the one hand and B on the other. This admits of two interpretations: either manganese atoms go into D positions, or the aluminium atoms are not completely sorted out into B positions, a minority of the aluminium atoms being found in D positions.

The second alternative would be analogous to the iron aluminium alloys\* containing a little more than the correct amount of aluminium to satisfy the formula  $\text{Fe}_3\text{Al}$ . Here most of the aluminium atoms occupy position B, but a certain proportion move over to position D, leaving B without its full complement of aluminium atoms.

To distinguish between the two alternatives, it is necessary to make use of the fact that the intensities of the two sets of superlattice lines differ considerably with different radiations (see Table III). This could not be explained by a partial distribution of aluminium between positions B and D. It can, however, be explained by placing manganese atoms in D and aluminium atoms in B. This distribution gives structure factors  $2\text{Cu} - (\text{Mn} + \text{Al})$  and  $(\text{Mn} - \text{Al})$  for types (2) and (3) respectively. The difference is  $2(\text{Cu} - \text{Mn})$ . As we have shown in a previous paragraph, the scattering power of copper

\* Bradley and Jay, 'Proc. Roy. Soc. A', vol. 136, p. 210 (1932).

Table VI—Comparison of Observed and Calculated Intensities for Three Different Structures

In order to make the observed intensities comparable with the calculated values, they must be reduced to the same scale. This process is complicated because the observed values are found at room temperature, whereas the calculated values apply only at the absolute zero of temperature. This causes the observed intensities to fall off too quickly with increasing values of  $\sin \theta/\lambda$ . The observed values should be related to the calculated values in the following way

$$I_{\text{obs.}} = I_{\text{calc.}} K e^{-\frac{B \sin^2 \theta}{\lambda^2}},$$

where  $\lambda$  is the wave-length and  $\theta$  the glancing angle.  $B$  is a physical constant which is the same for all radiations.  $K$  is arbitrary and differs for each experiment, depending on the experimental conditions. The scales of the observed values are adjusted by choosing values of  $K$  and  $B$  to give the best measure of agreement between the observed and calculated reflections for planes 220, 400, 422, and 440, the values of which are independent of the atomic arrangement. It was found that the best agreement between observed and calculated values was obtained by putting  $B = 3$ .

Table VI includes results from iron and copper radiations. The zinc values are not included as the difference in the scattering powers of copper and manganese is too small to give decisive results. It can be seen from this table that almost all the observed intensity values of the superlattice lines lie between the calculated values for structure (i) and structure (ii). Structure (iii) is, therefore, ruled out.

The obvious interpretation of this result is that structure (i) is essentially correct, but since there are too few manganese atoms to fill the whole of the D positions some of the manganese has been replaced by copper. This has the effect of reducing the difference in scattering power between A and C on the one hand and D on the other. The formula for the alloy may be written as  $\text{Cu}_2(\text{MnCu})\text{Al}$ . It must, however, be taken into account that there is a small proportion of aluminium atoms in excess. This must either replace copper in A and C or manganese in D. Good agreement with the observed intensities can only be obtained if it is supposed that the excess of aluminium is equally distributed between positions A, C, and D. The most probable atomic distribution is as follows —

A atoms	0.95 copper, 0.05 aluminium
B atoms	All aluminium.
C atoms	0.95 copper, 0.05 aluminium
D atoms	0.3 copper, 0.65 manganese, 0.05 aluminium

The calculated intensities for this structure are compared with the observed values in Table VII. The scale has again been fixed to give the best agreement for the strong reflections, which depend only on the composition and not on the manner in which the atoms of different kinds are distributed. The intensities of the superlattice lines are thus obtained on an absolute scale, and the close agreement with the calculated values can therefore be taken as a conclusive proof that both the structure is correct and also that the empirical rule for calculating  $f$  values in the neighbourhood of the K absorption edge is reliable.

Table VII—Comparison of the Observed and Calculated Intensities for the Most Probable Structure

Line		Iron $K_{\alpha}$ radiation $\lambda = 1.934$		Copper $K_{\alpha}$ radiation $\lambda = 1.539$		Zinc $K_{\alpha}$ radiation $\lambda = 1.434$	
$2\theta^{\circ}$	$hkl$	Observed	Calculated	Observed	Calculated	Observed	Calculated
3	111	15	15	32.5	30	37	40
4	200	37.5	37	38.5	39	39	38
8	220	353	364	587	589	655	647
11	311	5	5	11	11.5	10	11
13	222	10	8.5	8	9	7	8
16	400	59	57	86	87	95	95
19	311	1.5	2	4	4	3	4
20	420	12	12	9.5	9.5	6	7
24	422	161	153	169	164	172	171
27	{ 511 333 }	{ 2 2 }	{ 2 2 }	{ 2.5 3 }	{ 3 3 }	{ 4 4 }	{ 3 3 }
32	440	101	98	50	52	49	50
35	531	3.5	5.5	3.5	3	—	—
36	{ 600 442 }	{ 30 30 }	{ 32 32 }	{ 4.5 3.5 }	{ 3.5 3.5 }	{ — — }	{ — — }

In conclusion, the authors thank Professor W. L. Bragg, F.R.S., for his kind interest in the work, which was carried out in the Physical Laboratories of the University of Manchester.

### Summary

In an investigation of the ferromagnetic alloys of copper, manganese, and aluminium an alloy was found which showed an almost complete change of crystal structure due to heat treatment. Drillings of this alloy, which had been annealed at  $500^{\circ}$  for several hours and cooled slowly to room temperature, were found to have the  $\delta$  copper aluminium ( $\text{Cu}_3\text{Al}_4$ ) type of structure. The formula may be written as  $(\text{CuMn})_3\text{Al}_4$ , the atoms occupying the same positions as in  $\text{Cu}_3\text{Al}_4$ . The annealed and slowly cooled alloy is non-magnetic, but on

quenching from 800° C it becomes strongly ferromagnetic. The structure is now entirely body-centred cubic, with a face-centred superlattice.

A quantitative X-ray examination of the ferromagnetic alloy showed the utility of a new method for distinguishing between elements of almost equal scattering power. On comparing X-ray powder photographs of the same specimen made with radiations from iron, copper, and zinc anticathodes, it was found that the relative intensities of the weaker reflections varied with the wave-length of the radiation. This fact made it possible to distinguish the manganese atoms from the copper atoms, which would not have been possible if results from only one radiation had been available. The difference between the atomic scattering factors of copper and manganese which is very small with zinc radiation becomes fairly large with iron radiation.

The ideal structure of the ferromagnetic alloy would be —

Cu	0 0 0	0 $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ 0 $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ 0
	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ 0 0	0 $\frac{1}{2}$ 0	0 0 $\frac{1}{2}$
Al	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$
Mn	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$	$\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$

In the specimen examined there was a deficit of manganese atoms. The alloy was nevertheless homogeneous, some copper taking the place of the missing manganese atoms. The aluminium atoms in excess were distributed equally in the positions normally occupied by copper and manganese.



### *The Diffraction of Electrons in the Halogens*

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#### *Introduction*

Although the complete theory of the scattering of electrons by gas atoms must take into account the distortion of the incident and scattered waves by the atomic field, the exchange of electrons between the atom and the incident beam, and the disturbance of the atomic wave functions by the incident and scattered waves, a satisfactory explanation of the diffraction effects observed in the angular distribution of the elastically scattered electrons is obtained simply by considering the distortion of the incident wave by the undisturbed field of the atom\*. The scattering at large angles will then mainly depend upon the nature of the atomic field at the point in the atom where the potential energy of the incident electrons is equal to their kinetic energy.

Now the magnitude and gradient of the field at any point within the atom at a distance  $r$  from the centre is determined mainly by the nuclear charge and the screening constants of the electrons within the radius  $r$ , and hence the nature of the field at a point well within the outer electron shell will be similar for atoms whose electronic structures differ only in the constitution of the outer shell.

Thus, if the energy of the incident electrons is large compared to the ionization energy of the outer electron shell, we should expect that those elements of adjacent atomic number, which differ only in the structure of the outer shell, such as a rare gas and the adjacent alkali metal or halogen, would give similar diffraction patterns.

Some experimental evidence for this was recently obtained by Mohr and Nicoll† who showed that the angular scattering curves for argon, sulphur and phosphorus were very similar. Further evidence is provided by some previous results of mine‡ for  $\text{CH}_4$ ,  $\text{N}_2$ ,  $\text{CO}$  and  $\text{Ne}$ , the diffraction patterns of which are all very similar, as one would expect, since the scattering in these gases is

\* Except for very slow electrons and for the lightest gases when exchange effects become important.

† 'Proc Roy Soc,' A, vol 138, p 469 (1932)

‡ Arnot, 'Proc Roy Soc,' A, vol. 123, p 615 (1931)

mainly due to the C, N, O and Ne atoms which all occupy the same row of the Periodic Table and differ only in the structure of the outer electron shell

In this paper I present some results for iodine, bromine, bromoform and carbon tetrachloride. It will be seen that these results are very similar to those which I obtained some time ago for the rare gases \*

### *Apparatus*

The apparatus used in this work, and the experimental procedure adopted, have been fully described in previous papers †. It consists essentially of an electron gun producing a narrow homogeneous beam of electrons which can be set at different angles to a system of collecting slits, between which potentials are placed so as to separate the elastically scattered electrons from those which have excited or ionized the atom and from the positive ions so produced.

The apparatus was thoroughly baked out, the filament flashed and then run for about 15 minutes until the pressure read on a McLeod gauge fell to about  $10^{-5}$  mm of Hg. Mercury vapour was excluded from the apparatus by having a trap on the outlet tube immersed in a mixture of solid  $\text{CO}_2$  and alcohol. During a run the pumps were kept going and the vapour of the various substances was allowed to flow continuously through the apparatus, being admitted through an adjustable leak. The vapour pressure was measured by a Pirani gauge situated in the scattering chamber itself. The Pirani gauge was calibrated against the McLeod gauge with air, and it was then assumed that the calibration constant so obtained was approximately correct for the vapours used. During a run the pressure was kept constant at a value between  $1 \times 10^{-3}$  and  $3 \times 10^{-3}$  mm, of Hg. Over this range of pressure the relation between the scattered current and the pressure is known from previous work\* to be linear for argon. The linear relationship was again tested in this work with  $\text{CCl}_4$  and with  $\text{CHBr}_3$  and was found to hold, consequently it may be concluded that the results presented here are due to single scattering.

The substance in the vapour of which the scattering was to be measured was contained in a small bulb which communicated with the apparatus through a tap and the adjustable leak.

*Iodine*—About a gram of solid iodine was placed in the bulb which was then surrounded with cotton wool and kept at room temperature. The vapour pressure behind the leak was thus about 0.15 mm.

\* Arnot, 'Proc. Roy. Soc.' A, vol. 133, p. 615 (1931).

† Arnot, 'Proc. Roy. Soc.' A, vol. 129, p. 361 (1930), vol. 130, p. 655, vol. 133, p. 615 (1931), vol. 140, p. 334 (1933).

*Bromine*—The bulb containing about 1 c.c. of bromine was immersed in alcohol at a temperature of about  $-60^{\circ}\text{C}$  contained in a vacuum flask. The temperature of the alcohol was reduced by adding solid  $\text{CO}_2$  until the vapour pressure behind the leak was about 0.3 mm. The bromine was supplied by The British Drug Houses, Ltd., and was of their A.R. standard.

*Bromoform*—About 1 c.c. of bromoform was placed in the bulb which was then immersed in a mixture of crushed ice and water giving a vapour pressure behind the leak of about 2 mm. The bromoform was supplied by The British Drug Houses, Ltd., as free from alcohol.

*Carbon Tetrachloride*—The bulb containing a few cubic centimetres of  $\text{CCl}_4$  and a lump of anhydrous calcium chloride was surrounded with cotton wool and kept at room temperature. The vapour pressure in the bulb was thus about 70 mm.

The iodine and bromoform were dried by inserting a small glass tube containing  $\text{P}_2\text{O}_5$  into the bulb containing the substance with its open end above the surface of the substance. No drying agent was used with bromine as the vapour pressure of ice at  $-60^{\circ}\text{C}$  is negligible compared with the vapour pressure of bromine in the bulb. In each experiment during the evacuation of the bulb the substance was frozen with solid  $\text{CO}_2$  and alcohol.

After completing the work on one substance the apparatus was dismantled and thoroughly cleaned except after doing bromoform, which was the second last vapour to be used. When the work on bromoform was completed the bromoform was replaced by bromine, and the apparatus baked out at  $400^{\circ}\text{C}$  for about 2 hours. After the apparatus had cooled down, and with the filament on, the vapour pressure read on the Pirani gauge was less than  $10^{-4}$  mm. The reason for not dismantling the apparatus (and for not renewing the filament) was so as to have the conditions for scattering in bromoform and bromine as identical as possible, so that the magnitude of the scattering in  $\text{CHBr}_3$  and  $\text{Br}_2$  might be truly comparable.

#### *Discussion of Results*

The results, which have all been reduced to a common pressure and a common value for the intensity of the electron beam, are shown graphically in figs 1 and 2, and the experimental values used in constructing these curves are given in Tables I and II. As the results for all electron energies in any one vapour, except carbon tetrachloride, were obtained in a single run on the same day, the values for different energies in the same vapour should be strictly comparable. Owing to the precautions taken not to disturb the alignment of



alits and filament between the experiments with bromoform and bromine (see above) the values for these two vapours should also be strictly comparable.

Table I—Carbon Tetrachloride and Iodine

Carbon tetrachloride (CCl <sub>4</sub> )							Iodine (I <sub>2</sub> )				
V <sub>0</sub>	15	30	42	50	84	100	V <sub>0</sub>	15	42	80	122
θ°	I <sub>0</sub>	I <sub>0</sub>	I <sub>0</sub>	I <sub>0</sub>	I <sub>0</sub>	I <sub>0</sub>	θ°	I <sub>0</sub>	I <sub>0</sub>	I <sub>0</sub>	I <sub>0</sub>
25	224	103	—	112	—	76 0	18	—	—	84 0	93 0
30	—	—	113	—	60 0	—	23	—	—	27 5	38 0
35	98 0	76 0	—	50 0	—	25 3	28	—	52 1	18 0	34 0
40	—	—	78 5	—	24 9	—	33	—	30 5	21 5	41 7
45	79 0	57 0	—	20 5	—	11 6	38	274	16 8	26 1	37 9
50	—	—	23 7	—	10 9	—	43	—	11 4	23 4	30 3
55	66 5	25 5	—	9 4	—	0 15	48	155	9 45	17 7	22 5
60	—	—	12 7	—	7 23	—	53	—	9 65	—	10 6
65	44 0	12 8	—	7 1	—	0 20	58	108	10 8	6 85	4 38
70	—	—	10 5	—	5 85	—	63	—	12 1	—	—
75	31 4	12 0	—	7 9	—	5 40	68	68 2	12 0	2 57	4 70
80	—	—	10 0	—	5 60	—	73	—	10 9	—	—
85	31 2	15 1	—	11 3	—	6 54	78	51 7	9 15	5 41	15 3
90	—	—	16 4	—	6 40	—	—	—	—	—	—
95	33 4	19 6	—	12 0	—	5 65	88	40 7	7 50	6 25	18 1
100	—	—	16 8	—	6 00	—	98	35 6	10 6	5 30	7 22
105	32 3	20 7	—	13 0	—	4 82	108	35 3	20 0	6 90	3 23
110	—	—	15 0	—	5 45	—	118	35 6	26 0	10 6	3 40
115	31 6	17 2	—	11 3	—	4 30	128	35 0	22 6	9 90	6 60
120	—	—	13 6	—	4 87	—	138	32 6	14 2	4 99	4 73
125	33 1	12 8	—	10 2	—	0 00	148	29 2	11 5	5 35	5 08
130	—	—	11 0	—	5 62	—	—	—	—	—	—
135	39 0	11 6	—	10 0	—	9 80	—	—	—	—	—
140	—	12 1	12 3	11 0	8 62	—	—	—	—	—	—
145	—	—	—	—	—	15 6	—	—	—	—	—
150	—	13 7	14 3	15 3	14 7	—	—	—	—	—	—

The curves in figs 1 and 2 have all been displaced in a vertical direction, the ordinate zero for each curve being shown by the short horizontal lines at the side of the curves. The curves for 15- and 30-volt electrons in CCl<sub>4</sub> have not been displaced relative to each other, and consequently have a common base line. The ordinates of the 15-volt curve in iodine, bromine and bromoform have all been reduced by half. The number at the side of each curve represents the energy V<sub>0</sub> of the incident electrons in volts.

In fig 1 some curves for xenon and krypton, taken from an earlier paper,\* have been given for comparison. The ordinate values of the xenon curves (which are comparable for the different energies) have been adjusted so that the

\* Arnot, 'Proc Roy Soc,' A, vol. 133, p. 615 (1931).

maximum at 90° in the 122-volt curve fits the corresponding maximum in the 122-volt curve for iodine. The krypton values, which are also comparable, have been adjusted so that the maximum at 70° in the 80-volt curve fits the corresponding maximum in the 80-volt curve for bromine.

Table II —Bromoform and Bromine

Bromoform (CHBr <sub>3</sub> )					Bromine (Br <sub>2</sub> )				
V <sub>0</sub>	15	41	80	121	V <sub>0</sub>	15	41	80	121
θ°	I <sub>θ</sub>	I <sub>θ</sub>	I <sub>θ</sub>	I <sub>θ</sub>	θ°	I <sub>θ</sub>	I <sub>θ</sub>	I <sub>θ</sub>	I <sub>θ</sub>
24	—	—	—	146	24	—	—	144	82.0
29	485	253	110.5	74.0	29	485	220	78.0	36.8
34	—	108	65.5	43.7	34	—	138	34.2	16.6
39	284	129	74.8	30.1	39	285	79.0	16.4	10.4
44	—	73.5	24.5	24.4	44	—	53.1	9.52	8.97
49	221	51.0	22.9	22.5	49	182	30.8	8.77	9.45
54	—	37.9	22.8	20.5	54	—	23.0	10.4	10.2
59	167	28.3	22.7	19.3	59	121	19.6	11.7	10.3
64	—	25.5	19.0	17.2	64	—	19.4	12.2	9.80
69	117	23.5	16.3	13.2	69	90.0	20.1	12.1	7.82
74	—	22.5	14.7	—	74	—	19.3	11.0	6.34
79	76.8	21.0	12.3	6.80	79	75.0	17.7	9.62	3.84
84	—	19.3	—	—	84	—	16.2	—	—
89	66.1	16.8	7.62	4.65	89	65.8	15.0	5.40	—
99	62.5	14.0	6.90	8.02	99	61.4	15.7	2.87	4.00
109	68.2	13.9	9.28	12.0	109	58.7	18.2	3.80	6.90
119	78.0	17.5	12.5	13.2	119	59.0	21.6	5.57	7.96
129	85.7	23.3	15.2	10.3	129	63.4	21.1	6.00	6.30
139	96.7	31.0	15.6	8.59	139	71.5	19.7	4.99	3.64
149	—	28.2	12.6	6.65	149	—	19.9	4.16	2.62

In fig. 2 my argon results for 42-, 83- and 124-volt electrons have been reduced so that the maximum at 105° in the 42-volt curve fits the corresponding maximum in the 42-volt curve for CCl<sub>4</sub>. As my results extended only to 120° I have used the results of Mohr and Nicoll\* to extend these curves to 155°. The 30-volt curve for argon and the four curves for H<sub>2</sub>S are also due to Mohr and Nicoll. These five curves have all been fitted separately to the CCl<sub>4</sub> curves at the maxima.

\* 'Proc. Roy. Soc., A, vol. 133, p. 469 (1932). I have not used their complete curves because their values for different energies are not comparable. The extension of the 124-volt curve is not very accurate as it was obtained by interpolation between the ordinate values of Mohr and Nicoll's 160-volt curve and Hughes and McMillan's 100-volt curve ('Phys. Rev.', vol. 30, p. 585 (1932)).

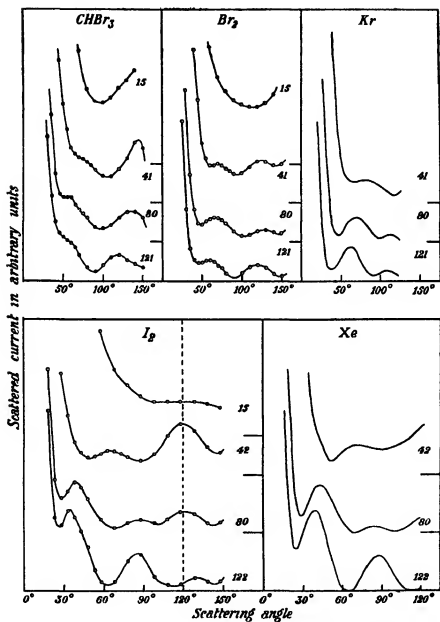


FIG. 1—Angular distribution of elastically scattered electrons in bromoform, bromine and iodine with curves for krypton and xenon given for comparison

Referring now to the curves for iodine and xenon, and remembering that the xenon curves extend only to  $120^\circ$  (i.e., up to the broken line in the iodine figure), we notice the remarkable similarity that exists not only in the shape of the curves but also in the relative magnitude of the scattering at the different energies. The various maxima and minima occur at approximately the same

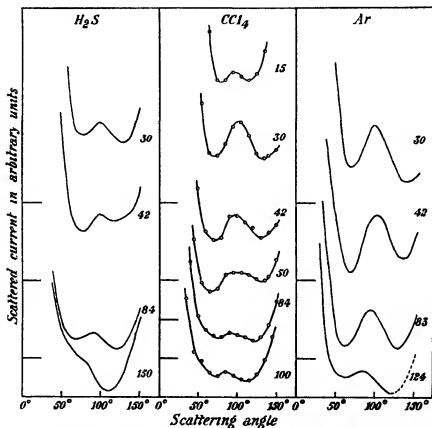


FIG. 2—Angular distribution of elastically scattered electrons in carbon tetrachloride, with curves for hydrogen sulphide and argon given for comparison

angle in both sets of curves, but the first minimum in the iodine curves is not quite so pronounced as it is in the xenon curves

The same general similarity is shown by the bromine and krypton curves, but it is not so marked as for those of iodine and xenon

The vapour of iodine and bromine at room temperature consists of diatomic molecules, but Massey and Mohr\* have shown that the scattering produced

\* 'Proc Roy Soc,' A, vol 135, p 258 (1932)

by a homonuclear diatomic molecule is practically the same as that caused by the two separate atoms. These results, therefore, provide further evidence that the scattering produced by atoms that differ only in the structure of the outer electron shell is practically identical for electrons having an energy greater than the ionization potential of the outer shell.

It is interesting to notice the remarkable similarity of the bromine curve for 41-volts and the iodine curve for 42-volts. In both curves there are maxima at  $70^\circ$  and  $120^\circ$ , the latter being higher than the former. The interesting and important point is that these curves in bromine and iodine are so very similar, whereas the two corresponding curves (40-volt curves) in krypton and xenon are not nearly so similar, neither are the curves for higher energies in bromine and iodine or in krypton and xenon so similar.

In view of the recent work of Childs and Massey\* it might be expected that curves for the same energy in krypton and xenon, or in atomic bromine and iodine would be similar, for they have shown that cadmium gives curves very similar to those found for mercury, which lies in the same column of the Periodic Table. They explain this by showing that the outer field of the cadmium atom is similar to that of mercury, since the two outermost shells in the two atoms have respectively the same number of electrons and approximately the same ionization potentials, and the same radii.

Table III

Shell	Sub shell	Xenon		Krypton	
		Ionization potential in volts	Radius in A	Ionization potential in volts	Radius in A
K	$1s^2$	33500	0.010	14300	0.015
L	$(2s, 2p)^8$	4800	0.041	1800	0.087
M	$(3s, 3p)^8$	647	0.112	160	0.194
	$3d^{10}$	518	0.146	74.5	0.324
N	$(4s, 4p)^8$	150	0.280	13.94	0.885
	$4d^{10}$	27.4	0.400		
O	$(5s, 5p)^8$	12.08	1.04		

This, however, is not so for xenon and krypton. In Table III are given the ionization potentials and the radii (defined as the distance from the centre at which the charge density of the shell electrons is a maximum) for the various

\* 'Proc. Roy. Soc.,' A, vol. 141, p. 473 (1933)

shells in xenon and krypton. The number of electrons in each shell is denoted by the superscript on the symbol giving the total and azimuthal quantum numbers of the shell. The values (except the ionizations potential of the outer shell for which the experimental value is used) have been calculated by Slater's rules\*.

It will be seen from the table that the 42-volt electrons in xenon will be scattered mainly by the field at about the centre of the N shell of 18 electrons. In krypton the 41-volt electrons will be scattered by the field between the N shell of 8 electrons and the M shell of 18 electrons. We should therefore not expect these curves in xenon and krypton to be very similar. The same will, of course, apply to scattering by *atomic* iodine and bromine since the fields of these atoms will be much the same as those for xenon and krypton.

In *diatomic* iodine and bromine, however, the ionization potentials of the outer shells will be different from those in the atom owing to the redistribution of the outer electrons. Owing to the striking similarity of the 40-volt curves in iodine and bromine the portion of the field that is mainly effective in scattering electrons of this energy must be very similar in nature in the two molecules—far more similar than it is in the atoms.

The energies of the inner shells will not be very different from those of the atom, and hence the scattering curves for fast electrons will be practically the same as those for the atom.

Considering now the curves for bromine and bromoform we see that the curves for corresponding energies are on the whole of a similar nature, the main difference being that the first maximum in the  $\text{CHBr}_3$  curves is not so pronounced as in the  $\text{Br}_2$  curves, while the minimum at about  $90^\circ$  is more marked in the  $\text{CHBr}_3$  curves.

The theory of the scattering by a molecule of  $\text{CHBr}_3$  is complicated by the fact that since the molecule has a permanent dipole moment the scattering will depend upon the orientation of the axis of the molecular field relative to the incident and scattered waves†. However, the general similarity of the curves given here for  $\text{CHBr}_3$  and  $\text{Br}_2$  shows that the scattering in  $\text{CHBr}_3$  is mainly due to the relatively heavy bromine atoms constituting most of the outer portion of the molecule.

\* 'Phys. Rev.', vol 36, p 57 (1930)

† When the molecule has no permanent dipole moment the treatment is considerably simplified for we can then consider the axis of the molecules to be orientated in random directions. This allows us to average the scattering over all orientations to obtain the experimental conditions (see Massey, 'Proc. Roy. Soc.' A, vol 129, p. 616 (1930)).

We can explain the increased prominence of the minimum at  $90^\circ$  in the curves for  $\text{CHBr}_3$ , as being due to a contribution from the carbon atom to the total scattering, for the field of the central carbon atom will be fairly exposed over the portion that is screened only by the light hydrogen atom. If we "uncover" more of the field of the central carbon atom by replacing the heavy bromine atoms by relatively light hydrogen atoms we obtain a scattering curve consisting solely of a minimum at  $90^\circ$  as is shown by the results\* for  $\text{CH}_4$ .

It is clear from the above considerations that the scattering produced by a complicated molecule such as  $\text{CHBr}_3$ , which would be difficult to obtain theoretically, can be explained qualitatively by simply summing the contributions to the total scattering coming from the separate atoms of the molecule, assuming that these scatter independently and paying due regard to the fact that the field of an atom may be effectively screened on one or more sides by relatively heavier atoms. Thus it appears that a suggestion which I originally put forward some time ago\* in explaining the results obtained for the scattering of low energy electrons in methane, namely, that we might be able to examine the structure and field of complex molecules by analysis of the diffraction patterns exhibited in the angular scattering curves, may prove to be of definite value.

Since the scattering by bromoform is mainly due to the bromine atoms we should expect that the scattering produced by carbon tetrachloride would be even more truly representative of the scattering by chlorine, for not only has the  $\text{CCl}_4$  molecule no permanent dipole moment but also the field of the central carbon atom is now practically completely screened by the four chlorine atoms (assuming the tetrahedral structure of the molecule).

The results given in fig. 2 support this conclusion, for we see that the  $\text{CCl}_4$  curves are similar to those of argon, and are intermediate in form between those of argon and hydrogen sulphide, just as the position of chlorine in the Periodic Table is intermediate between argon and sulphur. The fact that the maximum at about  $90^\circ$  in the  $\text{CCl}_4$  curves is not appreciably depressed shows how effective is the screening of the carbon atom by the four chlorine atoms.

### General Discussion

The elastic scattering of electrons by a spherically symmetrical field is given by†

$$I(\theta) = \frac{1}{4k^2} \left| \sum_{l=0}^{\infty} (2l+1) [e^{2i\delta_l} - 1] P_l(\cos \theta) \right|^2, \quad (1)$$

\* Arnot, 'Proc Roy Soc.' A, vol 133, p 615 (1931)

† Mott and Massey, "The Theory of Atomic Collisions," chap 11

where  $I(\theta)$  is the scattered intensity at an angle  $\theta$ ,  $k = 2\pi/\lambda$ , where  $\lambda$  is the wave-length of the incident electrons, and the  $P_l(\cos \theta)$  are Legendre polynomials having the values —

$$P_0(\cos \theta) = 1$$

$$P_1(\cos \theta) = \cos \theta$$

$$P_2(\cos \theta) = \frac{1}{2}(3 \cos^2 \theta - 1), \text{ etc}$$

The form of these curves is shown in fig 3 (c)

The incident electrons are resolved into beams of different angular momentum  $\sqrt{l(l+1)} \hbar/2\pi$ . For each value of  $l$  a value of the wave function is found which is zero at the origin and has the asymptotic form

$$\sin(kr - \frac{1}{2}\pi l + \delta_l),$$

which becomes

$$\sin(kr - \frac{1}{2}\pi l),$$

in the absence of a scattering field

The phases  $\delta_l$  in (1) are thus the phase differences between the asymptotic value of the wave function and its value when there is no atom situated at the scattering centre for each of these beams of different angular momentum. These phases for the various values of  $l$  determine the form of the angular scattering curve.

Now  $\delta_l$  decreases monotonically as  $l$  increases, and the series in (1) converges more rapidly the lighter the atom and the lower the energy of the incident electrons. Thus to represent the scattering of low velocity electrons by light atoms we require only the first few phases, and when this is so the predominant harmonic in (1) is that for which the corresponding phase has a value most nearly equal to  $\pi/2$ . This is clearly seen by separating the complex expression (1) into its real and imaginary parts. We then obtain

$$I(\theta) = \frac{1}{4k^2} \left[ \sum_{l=0}^{\infty} (2l+1) (\cos 2\delta_l - 1) P_l \right]^2 + \frac{1}{4k^2} \left[ \sum_{l=0}^{\infty} (2l+1) \sin 2\delta_l P_l \right]^2$$

Thus if  $\delta_l \approx \frac{1}{2}\pi$  then the angular scattering curve will be of the form  $|P_l(\cos \theta)|^2$ . For example, for 30-volt electrons in argon the calculated phases are\*  $\delta_0 = 2\pi + 0.885$ ,  $\delta_1 = \pi + 1.689$ ,  $\delta_2 = 1.938$ ,  $\delta_3 = 0.374$ ,  $\delta_4 = 0.159$ , and the scattering curve closely resembles the curve  $|P_2|^2$ .

Now as the velocity is increased the number of phases which have an appreciable size and which must therefore be taken into account in (1) increases ;

\* Holtmark, 'Z. Physik,' vol 55, p 437 (1929). In Table II of his paper  $\delta_1$  should be replaced by  $\delta_2 - \pi$ .



and when there are a large number of phases effective it is difficult to see from the above formula which harmonic (if any) will mainly determine the shape of scattering curve, for the effect of the higher harmonics will be increased by the weight factor  $(2l + 1)$ . However, even when a large number of terms must be taken into account in (1) it can be shown that the form of the curve at large angles is still due to the lower order harmonics, for all the higher harmonics interfere to form a curve which falls off monotonically to a small value at large angles. To see this we write the equation (1) in the form

$$I(\theta) = \frac{1}{4k^2} \left| \sum (2l + 1) 2i\delta'_l P_l(\cos \theta) + \sum (2l + 1) [e^{2i\delta_l} - 1 - 2i\delta'_l] P_l(\cos \theta) \right|^2 \quad (2)$$

Now the first series on the right is just Born's first approximation to the exact formula (1) where  $\delta'_l$  are the Born phases which for small values are equal to the exact phases  $\delta_l$ .

It is well known that for all atomic fields the scattering given by Born's formula falls off monotonically\* to a small value at large angles, and therefore the shape of the scattering curve at large angles will be determined by the second series on the right for all velocities less than that for which the Born approximation is valid, and it is clear that this series will contain only the lower order harmonics for which the phases have a value greater than about 0.5

Thus for electrons of high energy  $\delta_l = \delta'_l \ll 1$  for all values of  $l$ . Then the term in square brackets in (2) is zero, and the scattering curve is given by the Born formula. As the energy of the incident electrons is decreased the phases increase,  $\delta_0$  first becoming appreciable, then  $\delta_1$ , then  $\delta_2$ , and so on, and consequently deviations from Born's formula will be noticeable at large angles through contributions coming in from the first few harmonics. Thus when the energy is decreased until  $\delta_0 \sim 1$  the term in square brackets becomes effective, and at large angles the scattering curve is of the form  $|P_0|^2$ . On further decreasing the energy  $\delta_0$  increases and passes through the value  $\pi/2$  after which its effect on the scattering becomes less marked, but now  $\delta_1$  is approaching the value  $\pi/2$  and so the scattering curve takes the form  $|P_1|^2$ .

For example, in helium, when the energy of the incident electrons is greater than 500 volts, all the phases are very much less than unity, and the scattering

\* The first series of harmonics when summed forms a smooth curve since for no value of  $l$  does  $\delta'_l$  become strictly zero owing to the fact that the field of an atom has no finite boundary.

curve is given accurately by the Born formula. When the energy is reduced to 340 volts the zero order phase increases to the value  $\delta_0 = 0.696$  which makes the term in square brackets in (2) appreciable and so gives the scattering curve at large angles the form  $|P_0|^2$ , i.e., the curve remains practically flat instead of falling off uniformly with angle. On further decreasing the energy,  $\delta_0 \rightarrow \pi$  but  $\delta_1 \rightarrow 0$  and consequently  $\delta_1$  never becomes great enough to give the scattering curve\* the form  $|P_1|^2$ . We might, however, expect to find the Born curve changing to the form  $|P_0|^2$  and then to the form  $|P_1|^2$  in a heavier atom, and this is found to be the case in neon, as an examination of the curves in fig. 2 of a previous paper† clearly shows. For argon we see from fig. 3 (a) of this paper that as the velocity is decreased the scattering curve, which is of the form  $|P_0|^2$  at 780 volts, changes to  $|P_1|^2$  at about 330 volts, and finally to  $|P_2|^2$  at about 80 volts.

Now in order that the scattering curve should pass from the Born curve successively through the forms  $|P_0|^2$ ,  $|P_1|^2$ ,  $|P_2|^2$ , etc., as the energy is decreased, the phases of these lower harmonics must not be too close together in magnitude, for if they are, the contributions coming from these lower harmonics will be comparable, and so the scattering curve will be the result of the superposition of several lower harmonics‡. For light atoms the lower order phases are probably sufficiently far apart in magnitude (over the range of energies up to that for which the Born approximation is valid) for the form of the curve to pass successively from one harmonic to the next, and the results for helium, neon and argon described above strongly support this view. But for heavy atoms such as mercury this may no longer be true, and the scattering curves will then be more complicated.

As the energy of the incident electrons is reduced from the value at which the Born approximation is valid all the lower order phases at first increase. We have seen how this accounts, with light atoms, for the scattering curve passing successively from the Born form through the forms  $|P_0|^2$ ,  $|P_1|^2$ ,  $|P_2|^2$ , etc. Now as the energy tends to zero all the phases do not continue to increase. In helium  $\delta_0$  increases to  $\pi$ , but all the other phases pass through a maximum and then tend to zero as  $k \rightarrow 0$ . In argon  $\delta_0 \rightarrow 3\pi$ ,  $\delta_1 \rightarrow 2\pi$  and the higher phases tend to zero. In krypton  $\delta_0 \rightarrow 4\pi$ ,  $\delta_1 \rightarrow 3\pi$ ,  $\delta_2 \rightarrow \pi$  and the others tend to zero, and in mercury  $\delta_2 \rightarrow 2\pi$  and the higher phases tend to zero. Therefore, as the energy is decreased from the value at which the Born

\* The minimum in the curve observed at lower energies is due to electron exchange.

† Arnot, 'Proc. Roy. Soc.' A, vol. 133, p. 615 (1931).

‡ I am indebted to Dr. C. B. O. Mohr for drawing my attention to this point.

approximation is valid we should expect the scattering curve to take the form of successively higher harmonics down to a certain energy, and then to reverse the order and pass back to the form of the lower harmonics

That this actually does occur is seen by an examination of the curves for argon\*. We have already seen above that the curve passes successively through the forms  $|P_0|^2$  at 780 volts,  $|P_1|^2$  at 330 volts, to  $|P_2|^2$  at 83 volts. It maintains the form  $|P_2|^2$  down to 30 volts at which value  $\delta_2$  is close to  $\pi/2$ . On further decreasing the energy the phase  $\delta_2$  goes to zero rather rapidly,† and the curve changes back towards the form  $|P_1|^2$ . At 10 volts the curve has almost precisely the same form as it had at 167 volts.

Now for any one atom the phases at first increase as the energy is decreased, and the same will occur if we keep the energy constant and increase the atomic number of the scattering atom. We might, therefore, obtain a series of curves by scattering electrons of a constant energy in various gases of increasing atomic number which show the same variation in form as curves obtained by scattering electrons of decreasing energy in a single gas. A series of such curves is shown in fig. 3. In fig. 3 (a) a set of scattering curves for electrons of decreasing energy in argon is shown, in fig. 3 (b) is given a series of scattering curves for 83-volt electrons in various gases‡ of increasing atomic number, and in fig. 3 (c) the form of the harmonics  $|P_0|^2$  to  $|P_5|^2$ .

We have already described the curves in fig. 3 (a), and have explained their change of form from  $|P_0|^2$  to  $|P_1|^2$  and finally to  $|P_2|^2$  as being due to the phases  $\delta_0$ ,  $\delta_1$  and  $\delta_2$  increasing and passing through the value  $\pi/2$  in turn as the energy of the electrons is decreased. The same reasoning applies to the curves in fig. 3 (b). As the atomic number of the gas is increased the phases  $\delta_0$ ,  $\delta_1$  and  $\delta_2$  increase and pass in turn through the value  $\pi/2$ , so giving the curves the form  $|P_0|^2$ ,  $|P_1|^2$ , and  $|P_2|^2$ .

The agreement between the two sets of curves is very striking, and this together with the internal consistency of each set provides strong evidence in favour of the fundamental correctness of the interpretation of scattering curves as given in this section. We see that at large angles the shape of a scattering curve for any energy up to the energy at which the Born approxi-

\* A set of curves for energies from 780 volts to 11 volts is given by Mott and Massey in "The Theory of Atomic Collisions," p. 135.

† Holtsmark, 'Z. Physik,' vol. 55, p. 437 (1929).

‡ The curves for sulphur, phosphorus and carbon are due to scattering in  $H_2S$ ,  $PH_3$  and  $CH_4$ . The  $H_2S$  and  $PH_3$  curves were obtained by Mohr and Nicoll (*loc. cit.*), the other curves in fig. 3 are the author's.

matron is valid depends upon the first few harmonics, and that in light atoms, where the phases of low order are well separated in magnitude, the form of the curve is given approximately over the whole of the above range of energies, by the harmonic whose associated phase has a value most nearly equal to  $\pi/2$ .

From figs 3 (a) and 3 (b) we see that the order of the phase which determines the shape of the scattering curve at first increases as the energy is decreased from the value at which the Born approximation holds, and that it also increases

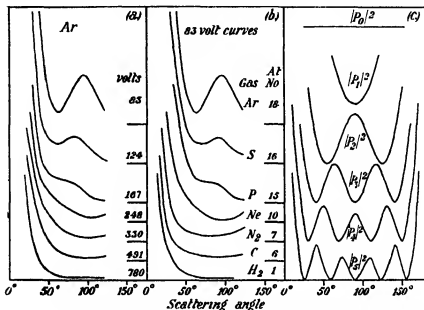


FIG. 3—(a) Scattering curves for electrons of various energies in argon (b) Scattering curves for 83 volt electrons in various gases of decreasing atomic number (c) Form of the curves  $|P_0|^2$  to  $|P_5|^2$

with the atomic number of the scattering atom. Thus the phase whose harmonic gives the form of the curve is the same for fast electrons in a heavy gas as it is for slow electrons in a light gas. This suggests that we may be able to determine the shape of the curve from the relation between the radii of the electron shells of the atom and a certain parameter  $r_0$  dependent upon the energy of the incident electrons. If  $r_0$  is defined as the distance from the centre of the scattering field to the point in the field at which the potential energy of the incident electron is equal to twice its initial kinetic energy, it is found that  $r_0$  must be approximately equal to the radius of the same shell for the scattering curve to have the same form in neon, argon and krypton.

The value of  $r_0$  can be found from the Hartree\* curves giving the effective nuclear charge for potential  $Z_p$  at various distances  $r$  from the centre of the atom. Tables of  $Z_p$  for various values of  $r$  are given by Holtmark† for argon and krypton, and by Brown‡ for neon. The values of  $r_0$  for various energies for which scattering curves have been obtained in krypton, argon and neon have been found from curves constructed from these tables. In fig 4 these values of  $r_0$  have been marked on the curves showing the radial distribution of

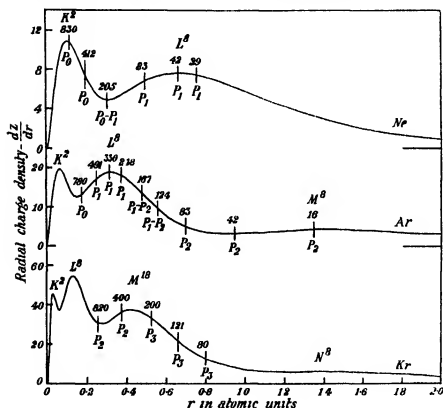


FIG 4—Showing, for electrons of various energies, the position of the incident electron with respect to the atomic shells when its kinetic energy is equal to half its potential energy in the atomic field

charge in the atoms. For argon and krypton I have used the charge distribution curves for the  $\text{Cl}^-$  ion and the  $\text{Rb}^+$  ion respectively given by Hartree,

\* 'Proc Camb Phil Soc,' vol 24, pp 80 and 111 (1928)

† 'Z Physik,' vol 55, p 437 (1929), vol 60, p 49 (1930)

‡ 'Phys Rev,' vol 44, p 214 (1933)

since Holtmark does not give values of  $-dZ/dr$ . These curves should be practically identical with those for argon and krypton.

The number at the top of the short vertical line marking the point  $r_0$  at which the kinetic energy of the incident electron is equal to half its potential energy in the atomic field, is the energy of the electron in volts, and below is marked the harmonic giving the approximate form of the scattering curve for electrons of this energy\*.

We see from fig. 4 that, for these three rare gases, when  $r_0$  is approximately equal to the radius of the K-shell the scattering curve has the form  $|P_0|^2$ , and hence it is  $\delta_0$  which most closely approximates to  $\pi/2$ . When  $r_0$  is equal to the radius of the L shell the curve has the form  $|P_1|^2$ , and when  $r_0$  is approximately equal to the radius of the M-shell the curve has the form  $|P_2|^2$  or  $|P_3|^2$ .

It also appears that we cannot expect the Born approximation to be valid at large angles unless the energy of the electrons is greater than the ionization potential of the K-shell. In mercury this potential is given by Slater's rules† as 75 600 volts. A rough extrapolation of the phase curves for mercury given by Heneberg‡ shows that  $\delta_0$  does not fall to the value  $\pi/2$  until the energy is increased to about 135,000 volts.

Calculations are in progress to determine the phases and theoretical scattering curves for the halogen atoms and molecules. The experimental work is also being continued in other molecular gases and vapours.

I wish to thank the Carnegie Trustees for a grant towards the cost of apparatus.

### *Summary*

The angular distribution of the elastically scattered electrons in the vapours of iodine, bromine, bromoform and carbon tetrachloride has been measured for several different energies of the incident electron beam. Diffraction effects are observed in all the curves. The curves for iodine, bromine and carbon tetrachloride are similar respectively to the curves previously obtained by the author for xenon, krypton and argon. The curves for bromoform show the same diffraction structure as do those for bromine.

These results, together with some previous results obtained by the author and other investigators, are then discussed from the theoretical standpoint.

\*  $P_1 - P_2$  means that the form of the scattering curve is intermediate between  $|P_1|^2$  and  $|P_2|^2$ .

† 'Phys. Rev.', vol. 36, p. 57 (1930).

‡ 'Z. Physik', vol. 83, p. 555 (1933).

Evidence is put forward to show that at large angles the shape of a scattering curve for any energy up to the energy at which the Born approximation is valid depends upon the first few harmonics, and that in light atoms, where the phases of low order are well separated in magnitude, the form of the curve is given approximately, over the whole of the above range of energies, by the harmonic whose associated phase has a value most nearly equal to  $\pi/2$

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### *On Metallic Dispersion in the Near Infra-Red*

By C. HURST, M.A., D.Phil., Christ Church, Oxford

(Communicated by F. A. Lindemann, F.R.S.—Received November 9, 1933)

Kronig\* has recently attempted to derive a quantum theory of dispersion in metallic conductors, using as a basis the Bloch theory of the metallic state, a more rigorous formulation of the same line of thought has been given by Fujioka†. In particular, Kronig has developed explicit formulæ for the coefficients of reflection and extinction and for the index of refraction of a metal in the so-called "transitional" region of the spectrum, where the period of the incident radiation is comparable with the mean time between two collisions of an electron with the metallic lattice. In order to obtain reasonable agreement with the room temperature measurements of Forsterling and Fréedericksz‡ in the near infra-red, it is necessary to assume a value for the electrical conductivity of the metal which is very much less than the value obtained by direct measurement. Kronig has suggested that this result may be due to the fact that the optical constants depend upon the properties of a thin surface layer of the metal, in which layer it is possible that the conductivity is diminished on account of boundary imperfections of the metallic lattice. It has already been shown, however, that even though this additional hypothesis be allowed, the formulæ in question are not consonant with the results of recent measurements§ on the high-temperature emissivities of metals in the near infra-red.

\* 'Proc Roy Soc,' A, vol 124, p 409 (1929), vol 133, p 255 (1931)

† 'Z Physik,' vol 76, p 537 (1932)

‡ 'Ann Physik,' vol 40, p 201 (1913)

§ Hurst, 'Proc Roy Soc,' A, vol 142, p 406 (1933)

The inadequacies of the quantum theory of the transitional region are more readily appreciated when it is realized that the formulæ for the optical constants derived therefrom are identical in scope with the formulæ of the dispersion theory of Drude,\* published in 1900. Kronig himself mentions that his formulæ tend, in the limit of zero frequency, to the classical, phenomenological relations which Drude† derived in 1894, but it does not appear to have been pointed out hitherto that the Kronig formulæ are, in fact, *identical* with, and not merely asymptotic to, those of the later dispersion theory of Drude.

In the latter theory, it was imagined that the free electrons in a metal were acted upon by a frictional force, proportional to their velocity, which represented the effect of their collisions with bound electrons and ions. If  $e$  and  $m$  are respectively the charge and mass of an electron, and  $N$  is the number of conduction electrons per unit volume, such an assumption leads to equations of the form

$$n^2 = k^2 - \Pi^2 - \frac{4\pi N e^2 m}{(2\pi m \nu)^2 + b^2}, \quad nk = \frac{b N e^2}{\nu \{(2\pi m \nu)^2 + b^2\}},$$

for the refractive index,  $n$ , of the metal and its absorption coefficient,  $k$ , at frequency  $\nu$ .  $b$  is the coefficient of the frictional force in the equations of motion of the free electrons, and is connected with the electrical conductivity,  $\sigma_0$ , of the metal by the relation

$$b = \frac{N e^2}{\sigma_0}$$

In the derivation of the formulæ, Drude assumed that the region of the spectrum under consideration is so far from the free periods of bound electrons and ions that their influence ( $\Pi^2$ ) on the index of refraction is constant and on the coefficient of absorption may be neglected.

Drude (*loc cit* 1904) himself came to the conclusion that this type of equation, with a constant value of  $b$ , would not suffice to represent the dispersion of metals, and pointed out that  $b$  might be expected to increase with the frequency,  $\nu$ , owing to stronger radiation losses at the higher frequencies. To explain the fact that the values of electrical conductivity given by the formulæ were too low, he, like Kronig, suggested a diminished conductivity of the surface layers of the metal. Forsterling and Fréedericksz compared their experimental results in the near infra-red with those given by the Drude equations, and failed to find agreement, they, too, put forward the hypothesis of an abnormally low surface conductivity.

\* 'Phys. Z.', vol 1, p 161 (1900). 'Ann. Physik,' vol 14, p 936 (1904).

† 'Physik des Aethers,' 1st Ed., p 574 (1894).



It is one of the great advantages of the modern conduction theory that no distinction need be made between free and bound electrons. Accordingly, Kronig (*loc cit*) has applied the Schrodinger dispersion theory to a metal lattice, regarded as a periodic field of potential, and has shown that, for sufficiently high frequencies, when the interaction of the electrons with the lattice may be neglected, the behaviour of an electron in a given stationary state,  $r$ , in an elementary cube of side,  $L$ , may be replaced, as far as its mathematical behaviour is concerned, by a fictitious system of harmonic oscillators, whose strengths are expressed in terms of the matrix elements of the linear momentum of the electron. As far as the near infra-red region is concerned, the most important oscillator (of strength  $f_r$ ) has a natural frequency zero. In other words, the spectrum of an electron,  $r$ , consists of the resonance frequency,  $\nu = 0$ , and a number of discrete absorption lines. Since the electrons in the metal are distributed over various stationary states, the equivalent spectrum of the electrons in the cube, in the absence of interaction, consists of the sharp resonance line and practically continuous absorption bands. In the visible region of the spectrum, the influence of the bands is predominant, while in the infra-red the resonance frequency is the more important. From about  $1.5 \mu$  onwards into the infra-red, the metal may be treated as though it had only the pseudo resonance frequency at  $\nu = 0$ , and the effect of the interaction of the electrons with the lattice may be introduced by discussing the broadening which this line undergoes thereby. Calling  $\delta$  the half-breadth of the representative line,  $\nu = 0$ , Kronig obtains the formulae

$$n^2 = \frac{\sigma_0}{\nu} \left( \frac{\delta}{\sqrt{\nu^2 + \delta^2}} - \frac{\nu\delta}{\nu^2 + \delta^2} \right), \quad k^2 = \frac{\sigma_0}{\nu} \left( \frac{\delta}{\sqrt{\nu^2 + \delta^2}} + \frac{\nu\delta}{\nu^2 + \delta^2} \right)$$

These equations are, however, identical in content with the Drude equations above (when the term  $NI^2$  is omitted), provided that

$$\delta = \frac{b}{2\pi m} = \frac{Ne^2}{2\pi m\sigma_0}, \quad (1)$$

or, since  $\delta$  is connected with the sum of the strengths,  $\sum_r f_r$ , of the fictitious resonators of frequency zero (where the summation is taken over all stationary states in which there are electrons) by the equation

$$\delta = \frac{e^2}{2\pi m\sigma_0 L^3} \sum_r f_r,$$

provided that

$$NI^2 = \sum_r f_r$$

Thus the only equation necessary for a complete correlation of the Drude and the Kronig dispersion formulæ is the trivial one expressing the fact that the zero frequency resonators are playing a rôle identical with that of the free electrons of the classical theory

That the results obtained by Kronig for the spectral range in question are precisely the same as those given by the classical treatment is not surprising as soon as it is realized that his method is closely analogous to that employed by Drude. Just as it is assumed in the older derivation of the formulæ that the influence of the free periods of the bound electrons and ions may be neglected, it is assumed in the quantum derivation that the contribution of the fictitious oscillators of non-zero frequency to the current induced in the metal by the incident light-wave is of little importance so long as the frequency under consideration is sufficiently low, the only excited electron states which are allowed to occur are therefore those which have slightly higher translational energies in the field-direction than usual. Again, when Kronig postulates that the effect of the interaction of the electrons with the metallic lattice is to cause a broadening of the remanent resonance-frequency, he is really making the assumption that the effect of the lattice vibrations is a shortening of the mean life-period of the excited electron states, and, as the electrons in excited states give their energy up to the lattice in the form of vibrational energy, which is ultimately distributed over all frequencies, the result is equivalent to the introduction of a damping factor, connected with the half-breadth of the resonance line by the relation (1)

Just as Drude regarded  $\sigma_0$  and his damping factor,  $b$ , as arbitrary parameters, so Kronig adjusts  $\sigma_0$  and  $\delta$  to make the values of  $n$  and  $k$  derived from the dispersion formulæ fit the experimentally observed values of Forsterling and Fréedericksz. As would be expected, the value of  $\delta$  computed for various wave-lengths for any one metal is not constant, but shows a tendency to decrease progressively with increasing wave-length, thus confirming the experience of Drude, already mentioned, concerning the behaviour of the damping factor,  $b$ . The comparison with experiment given by Kronig is, of course, superfluous, since Forsterling and Fréedericksz (*loc cit*) had already tested, in a slightly different manner, the same dispersion formulæ by means of their results.

As has so often occurred in the interpretation of other phenomena, the quantum-mechanical treatment of the problem of metallic dispersion in the near infra-red has proved to be nothing more than a new and rather more pleasing representation, in a different language, of classical concepts. Not

until an electron statistics which takes adequate account of the mutual potential energy of the electrons in a metal is developed, does there seem much hope of formulating a theory of the optical properties of metals in the near infra-red which will agree better than the classical theories with experimental data

In conclusion, I should like to thank Professor F A Lindemann, F R S, and Dr F London for their kind and helpful criticisms

### *Summary*

It is shown that the equations for the index of refraction and the coefficient of absorption of a metal in the near infra-red, which have been developed by Krong on a quantum-mechanical basis, are formally identical with the dispersion equations obtained by Drude in 1900 by classical methods. The close analogy between the underlying assumptions of the two derivations is indicated

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## *Photographs of Fluid Flow Revealed with an Ultramicroscope*

By A FAGE, A R C Sc

(Communicated by G I Taylor, F R S—Received November 14, 1933)

[PLATES 14-15]

1 *Introduction*—When an intensely bright beam of light from an ultramicroscope is passed through moving water, minute impurities which are always present and which cannot be seen in ordinary light, even with a microscope, become visible provided that they are seen against a dark background. These illuminated particles reveal the internal motions of the water without any interference with the flow, such as might be caused if extraneous particles or colouring matter were introduced into the water.

The ultramicroscope has recently been used to examine turbulent flow in pipes,\* and to obtain information on the flow in the wakes behind bluff obstacles†. An attempt has now been made to take photographs of some of the views seen when this earlier work was in progress. These photographs

\* Fage and Townend, 'Proc Roy Soc,' A, vol 135, p 656 (1932). Also 'Aero Res Ctee,' R & M 1474.

† Fage, 'Aero Res Ctee,' R. & M. 1510.

are given in the present paper\*. Photographs of the flow around a long circular cylinder at low Reynolds' number are also given. All the photographs, with the exception of those of fig 4 (c) and (d), Plate 14, are of water direct from the main. It is perhaps of interest to mention that the number of particles (probably mostly minute air bubbles) in the water appeared to differ appreciably from day to day.

2 Photomicrographs of the Brownian movements of ultramicroscopic particles in soap solutions have been taken by Darke, McBan, and Salmon†. These authors found that the finer details of the movements were not recorded. It appeared doubtful, therefore, whether satisfactory photomicrographs of moving water under a high magnification could be obtained, when, as in the present experiments, the particles in the water moved much faster than those which show the Brownian movement in a stationary fluid. Nevertheless, an attempt was made to take photographs with a Leitz photo-micrographic camera. Two makes of fast plate were used: Ilford Hypersensitive Panchromatic and Ilford Double X-Press Orthochromatic. A few streaks made by exceptionally bright particles were recorded, but details of the flow were not portrayed. The next steps possible were to increase the intensity of the light beam, or to decrease either the speed of flow or the magnification. The last of these steps was the most convenient to take, and it was decided to attempt to obtain photographs at a reduced magnification, and if these were satisfactory to enlarge the negatives. A small camera with an  $f/3$  lens ( $f = 2$  inches) and a Compur shutter was constructed. Photographs taken with this camera are given in figs 4-7, Plates 14, 15. The views were taken under a magnification of about 4 to 1 (except those for the prism, magnification 2 to 1) and the prints used for reproduction purposes were obtained from negatives enlarged 2.3 times. Ilford Hypersensitive Panchromatic and Ilford Double X-Press Orthochromatic plates ( $4\frac{1}{2} \times 6$  cm) were used, and they were found to give closely similar results. The exposure times varied from about 1 to 6 seconds.

3 *Turbulent Flow in a Pipe*—Photographs of turbulent flow in a square pipe of internal side 1.372 inches at a section situated 57.8 inches from the entry are given in fig 4. The mean rate of flow ( $U_0$ ) was about 0.7 ft per second. The corresponding value of  $U_0 m/\nu$ , where  $m$  is the hydraulic mean

\* The photographs were taken in the Aerodynamics Department of the National Physical Laboratory, and permission to communicate them was kindly granted by the Aeronautical Research Committee.

† 'Proc Roy Soc,' A, vol 98, p. 395 (1920-21)

depth, was about 1700. As described in the earlier work,\* the illuminated particles appeared as bright streaks inclined at various angles to the mean direction of flow, and these streaks appeared to intersect each other, owing to the persistence of vision. The maximum obliquity to the pipe axis of the streaks at a point in an axial plane XOY normal to a wall of the pipe increased slowly with the distance  $y$  from the axis, and reached a maximum at about  $y = 0.8s$ , where  $2s$  is the side of the pipe. The obliquity then fell to zero at

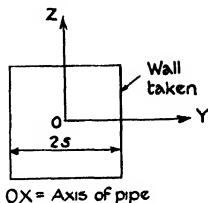


FIG 1

the wall ( $y = s$ ), so that particles viewed along and very close to the wall appeared to be moving in laminae parallel to the wall. Fig 4 (b) and (c) illustrate these features of the flow.

The maximum obliquity of particles viewed on a plane parallel to a wall increased continuously as the wall was approached, fig 4 (d). Cerebos salt was added to the water before taking photographs, fig 4 (c) and (d), to increase the number of bright particles near the wall, and so to improve the views.

The most interesting views seen in the earlier work were obtained when the microscope (magnification 200) was focused on the inner surface of a wall. Particles within a surface film of about 0.001 inch thick were then in focus. The view obtained showed all the particles to be moving in sinuous paths and occasionally paths of several wave-lengths were seen. It was also observed that the slowest particles, which were estimated to be about 1/40,000 of an inch from the surface, often moved in short jerks. An attempt was made to photograph with a Leitz photo-micrographic camera the details of this interesting motion, but without success. One of the difficulties experienced was that the intensity of light falling on the photographic plate from the illuminated

\* *Loc cit.*, § 1

particles was greatly reduced by the high magnification necessary to obtain the view. Moreover, it was found impossible to illuminate a very thin film without illuminating the surface itself and the light scattered from the surface tended to fog the plate. A photograph of laminar flow in the pipe is given in fig 4 (a).

4. *Triangular Prism*—Photographs of the turbulent wake behind a long triangular prism are given in fig 5 (a) and (b), Plate 14. These photographs illustrate the work described in the earlier paper (R & M, 1510), which was undertaken to show that at a sufficiently high Reynolds' number the eddying wake behind a very long cylindrical obstacle, mounted with its length normal to the stream eventually settles down to a regime in which the velocity disturbances are three-dimensional, even although the mean flow in planes at right angles to

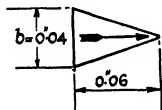


FIG 2

the length tends to be two-dimensional. The prism was mounted between the opposite sides of a 1.125 inch water tunnel. The dimensions of the triangular cross-section were 0.04 inch (base) and 0.06 inch (height). The photographs were taken at  $U_0 b/\nu = 250$  ( $b$  = base of section), at a point situated in the middle of the turbulent wake at a distance  $35b$  behind the prism. Fig 5 (a)

gives a view of the flow in a plane at right angles to the axis of the prism. It is seen that the disturbances are greatest over the central part of the wake, and that these disturbances gradually die away as the outer limits of the wake are approached. The maximum deviations of the streaks in the middle of the wake from the direction of mean flow were observed to be about  $\pm 17^\circ$ . Photograph 5 (b) was taken on the plane passing through the axis of the prism and shows that the flow is disturbed along the whole length of the prism. The maximum deviations observed in this plane were about  $\pm 23^\circ$ .

5. *Short Circular Cylinder*—Photographs of the flow at the exposed end of a circular cylinder extending from a wall to the centre of the tunnel are given in fig 6, Plate 14. The diameter,  $D$ , of the cylinder was 0.04 inch. These photographs were taken at a water speed,  $U_0$ , of 0.58 ft per second ( $U_0 D/\nu = 160$  approx.) Fig 6 (b) shows clearly the traces made by the axial flow of particles into the dead-water region behind the top of the cylinder.

6. *Long Circular Cylinder*—The photographs given in fig 7, Plate 15, show the changes with  $U_0 D/\nu$  in the flow around a long circular cylinder of diameter 0.04 inch. The cylinder was mounted with a small clearance of about 0.01 inch at one end between the walls of the water tunnel. The distance between the walls

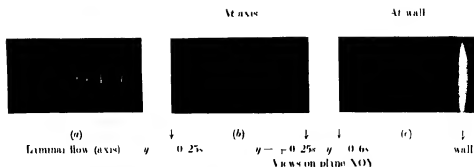


FIG. 4. Turbulent flow in pipe 1 cm.  $\nu = 1700$ .



FIG. 5. (a) Views on plane parallel to wall at  $y = 0.9x$  (about).



FIG. 5. Flow in the centre of the wake at a point 15*b* behind triangular prism.



FIG. 6. (a) Flow at the end of circular cylinder. (b) Flow at the end of circular cylinder.

Views at  $O$  on plane  $XOY$



(b)  
1 0 1 2  
v 17 7

(c) 21

(d) 27

(e) 32



(a) Free stream



(f) 47



(g) 104



(h) 170

(a)

Views at  $O$  on plane  $XOZ$



(i)



(j)



(k)

FIG. 7. Circular cylinder. Diameter = 0.04 m. length 1.12 m.



was 1.125 inches. Photograph (a) shows the steady nature of the flow in the empty tunnel. Photographs (b)-(h) were taken on the median plane XOY at right angles to the axis of the cylinder at values of  $U_0 D/\nu$  lying within the range 17.7 to 170. The flow patterns shown in these photographs of tap water resemble those obtained from experiments of other workers\* in which extraneous particles were introduced into the fluid (air and water). The first photograph (b) shows a flow which is obviously not far removed from the symmetrical laminar type. Fig. 7 (c) to (f) show the progressive changes in

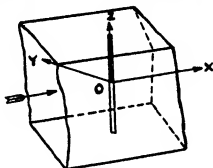


FIG. 3

the standing-vortex regime as the value of  $U_0 D/\nu$  is increased in steps over the range 21 to 47. The gradual increase in the length of the vortices with Reynolds' number will be noticed. Eventually, the standing vortices are drawn out to such a length that they break down and give place to a dead-water region behind the cylinder. Fig. 7, photographs f, g, and h illustrate the sequence in this breakdown.

Photographs i, j, and k were taken on the plane XOZ passing through the axis of the cylinder. Of these photographs j is the most interesting for it clearly shows that the flow in the standing vortices at  $U_0 D/\nu = 38.0$  is of a spiral character. This spiral motion is associated with an inward flow of particles from the ends of the cylinder. It will be noticed that the angles of the spirals become progressively smaller, and eventually zero, as the plane of symmetry XOY is approached.

7. In conclusion, the writer wishes to acknowledge his indebtedness to Miss V. M. Emms for assistance in the photographic work.

\* Nisi and Porter, 'Phil. Mag.' (1923). Also Nayler and Fraser, 'Aero. Res. Ctce.', R. & M. 332 (1917-18).

*Summary*

The ultramicroscope has recently been used to examine turbulent flow in pipes, and to obtain information on the eddying wake behind a bluff obstacle. Photographs are given in the paper of some of the views seen when this earlier work was in progress, also of the flow around a circular cylinder at low Reynolds' number. The flows were revealed by the motions of those minute particles in water which became visible in the light beam of the ultramicroscope.

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*The Thermal Decomposition of Nitrous Oxide at Pressures up to Forty Atmospheres*

By E HUNTER

(Communicated by C. N. Hinshelwood, F.R.S.—Received November 16, 1933)

Nitrous oxide decomposes to nitrogen and oxygen at velocities which can be conveniently measured at temperatures between 600° and 850° C. M. A. Hunter\* investigated the reaction by streaming the gas through a porcelain bulb in a furnace and measuring the decomposition for different times of passage. No attempt was made to determine whether the reaction is homogeneous or heterogeneous. The effect of wide variation of pressure was not used to determine its order, since the reaction was followed only over small ranges of decomposition at atmospheric pressure. From the velocity of decomposition, however, bimolecular constants were obtained which could be represented by the equation

$$\ln k = 24.12 - 31800/T,$$

where  $k$  is the bimolecular velocity constant and  $T$  the absolute temperature. If this equation holds, the activation energy of the bimolecular reaction is 62,040 cal./gm. mol.

A much more thorough examination of the reaction was made by Hinshelwood and Burk,† who measured the rate of reaction by following the pressure increase at constant volume in a silica bulb. The reaction was proved to be

\* 'Z. phys. Chem.', vol. 53, p. 441 (1905).

† 'Proc. Roy. Soc. A', vol. 106, p. 284 (1924).

homogeneous The initial pressure was varied between 50 and 500 mm Hg, and it was found that the reciprocal of the half-lives when plotted against the initial pressures gave a straight line True bimolecular reaction requires the straight line  $1/t_1 = ka$ , where  $t_1$  is the half-life, and  $k$  the velocity constant, and  $a$  the initial concentration The line through the experimental points showed a small intercept on the  $1/t_1$  axis for which no explanation was offered at the time From the variation of the bimolecular constants between 565° and 852° C the activation energy of the reaction was calculated to be 58,450 cal /gm mol If the reaction were a bimolecular one dependent on immediate decomposition at each activating collision of the molecules the number of molecules reacting per second should be equal to  $Z \times e^{-E/RT}$ , where  $Z$  is the number of molecules colliding per second and  $E$  is the activation energy From the observed rate of reaction at 1000° K a value of 55,000 cal /gm mol was found for the activation energy The fairly close agreement between the two values of the activation energy, 58,450 and 55,000 cal /gm mol and the manner in which the half-life varied with pressure provided good grounds for believing the reaction to be a simple bimolecular one, dependent only on collisions between the molecules

Further work on the reaction has been published by Volmer and Kummerow,\* who made velocity measurements at 665° C between 25 and 300 mm Hg They came to the conclusion that the velocity of reaction was simply that to be expected from the low pressure part of a quasi-unimolecular reaction of the type, which according to the theory of Landemann† may show a half-life independent of pressure over a very wide pressure range and yet exhibit a falling off of velocity at some sufficiently low pressure Volmer and Kummerow expected to find that the nitrous oxide decomposition would give true unimolecular constants at some sufficiently high pressure In a paper by Volmer and Nagasako‡ results at pressures up to 8000 mm Hg were given, and it was claimed that these agreed with the theory of a quasi-unimolecular reaction and showed that the unimolecular constants were practically independent of pressure above a pressure of about  $6\frac{1}{2}$  atms The activation energy was 53,000 cal /gm. mol

Some measurements made by Musgrave and Hinshelwood§ during a study of the reaction at low pressures are shown in fig 1 The line is almost straight

\* 'Z phys Chem,' B, vol 9, p 141 (1930)

† 'Trans Faraday Soc,' vol 17, p 598 (1922)

‡ 'Z phys Chem,' B, vol 10, p 414 (1930)

§ 'Proc Roy Soc,' A, vol 135, p 23 (1932)

near the origin, but passes through a region of great curvature at about 50 mm Hg and thereafter straightens out to a line of a much smaller slope than that near the origin. It is difficult to explain these results on the basis of Volmer's theory of a simple quasi-unimolecular reaction, since this would give a curve rising smoothly from the origin finally becoming horizontal, without the marked change of curvature and straight lines of fig. 1. Musgrave and Hinshelwood were led to the conclusion that the observed reaction was the sum of a unimolecular reaction, the line OAB of fig. 1, and a bimolecular reaction. The intercept OC' was regarded as a measure of the full extent of the unimolecular reaction and its variation with temperature gave a value of 50,500 cal/gm

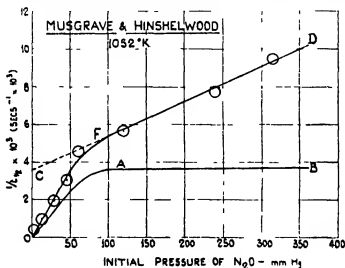


FIG. 1

mol for the activation energy of the unimolecular reaction. In the same paper Musgrave and Hinshelwood gave results which proved that the nitric oxide formed as a by-product catalysed the reaction slightly.

If Volmer's theory of a simple quasi-unimolecular reaction is correct, the half-life should be almost independent of pressure at all pressures above approximately 6 atms. On the other hand, Musgrave and Hinshelwood's theory of two concurrent reactions, one of which is bimolecular, suggests that the half-life might still be dependent upon pressure even above 6 atms. In order to discover the true state of affairs velocity measurements at pressures much greater than 6 atms are needed. Such results are provided by the present paper, which reports measurements of the velocity of decomposition of nitrous oxide at pressures as high as 40 atms.

## EXPERIMENTAL

*Apparatus*

The reaction was studied at temperatures between 840° and 999° C and at pressures between 0.10 and 40 atms, by measuring the rate of pressure increase of the gas at constant temperature and volume in a silica bulb. Two types of apparatus were used. The low pressure apparatus was designed for work at pressures between 50 and 2500 mm Hg, the high pressure apparatus for work at pressures up to at least 40 atms.

(1) *Low Pressure Apparatus*—The apparatus used by Hinshelwood and Burk (*loc cit*) and described by Hinshelwood and Topley\* was used for pressures up to 500 mm Hg and was adapted for use at pressures up to 2500 mm Hg by fitting spring-loaded taps to stand the pressure, and by employing a closed manometer for the pressure measurement. This manometer was made from a J-shaped glass capillary tube, 1 mm internal diameter. In the longer limb, dry CO<sub>2</sub>-free air was enclosed by a mercury thread, the movement of which in the shorter limb was used to indicate the pressure. The enclosed air was kept at a constant temperature by fitting the longer limb with a water-jacket through which water from a 25° C thermostat was circulated. The manometer was calibrated against an open mercury column and then sealed to the apparatus. The advantage of using a closed manometer of this type is that it entails only a small dead space (1.5 to 2% of the volume of the reaction bulb) and only a small correction to the results obtained with it is needed.

The nitrous oxide was stored in a glass bulb over mercury at a pressure of about 3000 mm Hg. At pressures up to 2500 mm the highest used with this apparatus, the gas appeared to reach the temperature of the reaction bulb within a second of its entry.

A change was made in the method of temperature measurement. In place of a millivoltmeter calibrated in degrees Centigrade, which was found to be liable to changing zero errors, the calibrated platinum-platinum rhodium thermocouple was used in conjunction with a potentiometer which made it possible to control the temperature to  $\pm 0.3^\circ \text{C}$ .

(2) *High Pressure Apparatus*—No apparatus for this type of work at pressures up to 40 atms has been described. Volmer and Nagasako (*loc cit*) used the static method at pressures up to 10 atms. They made little alteration to the apparatus used for normal pressures, the range of which is limited by the strength of the silica bulb.

\* 'J Chem Soc,' vol 125, p 393 (1924)

The solution of the present experimental problem seemed to be to have a steel bomb whose walls could be kept at a low, safe temperature, serving as a container for an electric furnace surrounding the reaction bulb. There remain the difficulties of keeping the reaction vessel at a uniform and accurately measurable temperature, of keeping down the dead space, and of providing a sufficiently sensitive means of pressure measurement. It is believed that these difficulties have been overcome in the apparatus described below, which was designed for use at pressures up to 100 atms.

*The Bomb, Furnace, and Reaction Vessel* A diagram of the steel pressure vessel and its contents is given in fig 2. The bomb A was made of mild steel, the cover plate B of "Vibra," and the two gave a pressure-tight fitting on the copper ring C by the tightening of eight nuts over the cover plate.

It was necessary to have insulated leads into the bomb for the current supplying the furnace and for the thermocouple. The chrome-steel glass-insulated cones\* which have been in use for some time at Winnington were used. For the furnace a single insulated cone was enough, the return being via the steel body of the apparatus. The copper wire leads to the furnace were threaded through a length of twin-bore "Vitreol" tube within the steel pressure tubing D. One wire was fastened to the inner terminal of the insulated cone E, the other was connected to the metal at F. By using the length of tube D, the cone was kept away from the heat of the furnace, which it was feared might have caused a breakage of the glass, if the cone had been fixed directly into

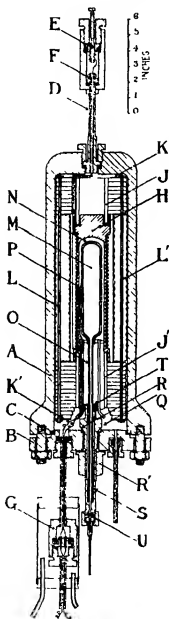


FIG 2

\* Welbergen, 'J Sci Instr., vol 10, p 247 (1933)

the body of the bomb. The outer part of the cone E was connected through rheostats to the positive lead of a 200-volt D.C. supply. The negative was connected to the steelwork of the apparatus. The glass insulation of the cones was much less than a millimetre thick, but care was taken to keep the bomb free from moisture, and it was found that the cone would carry a current of at least 3 amps. throughout the pressure range without trouble.

For the thermocouple leads into the bomb, two insulated cones set into one plug were needed. The platinum and platinum-rhodium wires from the hot junction were fixed to the inner terminals of the two cones, and from the outer terminals platinum and platinum-rhodium wires ran to cold junctions with copper, kept at the temperature of melting ice. The readings of a couple arranged in such a way are only correct if all the four junctions to chrome steel are at precisely the same temperature. Because of this it was not permissible to fix the pressure cones directly into the cover plate B, where they would have been subjected to a temperature gradient along their length, as a result of the dissipation of heat from the furnace. The pressure joint was accordingly made in the separate vessel G which was attached to the cover plate by a length of steel tubing along which passed the insulated wires from the hot junction. The vessel G was water-jacketed, so preventing the occurrence of any difference in temperature along or between the insulated cones.

A connection through the cover plate was made for the entry of the nitrogen used to balance the pressure within the reaction vessel.

The furnace was designed to give as even and as steady a temperature as possible in the heating space. It was rather a difficult problem because of the necessity of having a lagging which would take up a temperature gradient between the furnace wall at 600° to 900° C., and the inner bomb wall which it was hoped to keep below 200° C. It was found to be unavoidable that the furnace space would not be much greater than the required volume of a reaction bulb. This increased the difficulty of ensuring an even temperature throughout the furnace and reaction bulb, for with the bulb filling most of the furnace space it became important to minimize any tendency for the furnace temperature to fall away at the ends.

The tendency for a furnace to show this effect of a serious temperature gradient at its ends is usually due to the efficiency of the endwise lagging of the furnace being much poorer than that of the lagging of the sides. In the design of the furnace shown in fig. 2 special attention was given to this point. The nichrome heating wire, 70 ohms resistance, insulated with porcelain "fish spine" beads, 3 mm. in diameter, was wound in a deep helix cut in the

steel tube H. This metallic tube was used for the purpose of giving a good heat conductivity along the length of the furnace which would help to maintain an even temperature. The heating wire was covered with a thin layer of alundum cement, over which was wound sufficient asbestos string to give a diameter which would just fit the bomb. The asbestos was held in place by a jacket of wire-netting. The tube H was supported and prolonged by two lengths of wide "Vitreosil" tube J and J'. Over these were pressed a sufficient number of disks of asbestos cement board to occupy the space between the ends of the steel furnace tube and the ends of the bomb. The hollow space within the "Vitreosil" tube J was filled up with alundum cement. The whole furnace assembly was made rigid and self-supporting by bolting together two thin steel end-plates K and K', with two lengths of steel tube L and L' screwed at the ends. Tubing was used instead of rod to give a free passage for the gas between the front and the back of the bomb.

The reaction bulb M was made of silica and was about 16 cm. long with a volume of 110 c.c. To promote a rapid equalization of temperature over the whole surface of the silica, the bulb was covered with a closely fitting jacket N, made from sheet silver about  $\frac{3}{4}$  mm. thick. This jacket was made in two sections, one a test-tube shaped piece covering the whole length of the reaction vessel, the other a hemispherical cap which covered the remainder of the silica surface. The silica bulb was sealed to the silica capillary tube O, round which were arranged three "Vitreosil" tubes to pass gas into the furnace space, and a length of twin-bore "Vitreosil" tube to carry the thermocouple wires, over all these was wound asbestos string to the diameter of the tube J'. The hot junction of the thermocouple was fixed in contact with the mid point of the silver jacket at P, by looping it round a small tang of silver raised from the body of the jacket N. The leads to the junction were kept out of contact with the silver by threading them through 1 mm. bore "Pythagoras" refractory tubing.

The above type of construction gives a reaction bulb with an outer surface of a good heat conductor, heated in a furnace tube made of a good conductor, surrounded by a considerable thickness of lagging, particularly at the ends, so that the requirements for as even a temperature as possible are fulfilled.

One of the earliest difficulties was to find a way of making the joint between the reaction bulb and the steel gas supply pipe. The first attempts were made with butt joints, for which a flange formed on the end of the silica capillary O was held against a flange on the incoming steel tube, with a "Klingerte" washer between. The first joint made in this way was satisfactory,



but after a breakage many attempts to repair the joint were unsuccessful and the principle was discarded. The type of joint finally used was found to be quite satisfactory. It is shown in fig. 2. The mild steel tube Q was turned with a cone R, which gave a gas-tight fitting on the cover plate when the nut R' was tightened. It was found possible to join the silica to the steel by soldering the capillary O to the inside of the tube at a point S, sufficiently remote from the heat of the furnace to avoid softening the solder. From the solder at S to the interior of the bomb there was a clearance of about a millimetre between the silica and the steel. Some support was given to the capillary by the tightly fitting brass sleeve T. The length of the tube Q outside the bomb was cooled by a jet of air to make sure that there would be no softening of the joint S. The steel tube was connected to the gas supply and manometer system by a lens ring joint at U to 0.7 mm. bore copper capillary tube, running to a T-piece (i, fig. 3).

*Pressure Measurement*—The half-life, in the decomposition of nitrous oxide, is reached when the pressure is 1.25 times the initial pressure. For an accuracy of 1% a pressure measuring instrument sensitive to 0.25% of its pressure reading would be wanted. Closed gas manometers could not conveniently be used because of the difficulty of achieving this sensitivity and at the same time avoiding a large dead space. A pressure balance\* affords the most sensitive and accurate method of measuring pressures of the order of 100 atms., but it could not readily be adapted to follow the pressure change during reaction. It provides a reservoir of fluid at a constant pressure, which can only be changed by changing the load on the balance, an operation which takes an appreciable length of time, while the reaction pressure to be measured would rise smoothly and continuously from the instant of filling the bulb.

It seemed best to use sensitive carefully calibrated Bourdon tube gauges. Six gauges were obtained and calibrated against a pressure balance. Four of these were Schaeffer and Budenberg gauges with 12-inch dials, two for 100 kg/cm<sup>2</sup> and two for 50 kg/cm<sup>2</sup>. The remaining two gauges were Budenberg gauges for 300 lbs/sq. inch. With practice it was found possible to read these gauges to 0.05 kg/cm<sup>2</sup>, 0.02 kg/cm<sup>2</sup>, and 0.4 lbs/sq. inch respectively, so that they were sufficiently sensitive for the purpose for which they were required.

If these gauges had been used connected directly to the gas, the volume of the Bourdon tube, which was several cubic centimetres, would have been a very serious addition to the dead space of the apparatus. The presence of such a

\* Michels, 'Ann. Physik', vol. 72, p. 285 (1923), vol. 73, p. 577 (1924).

large dead space was avoided by filling completely with oil the gauge reading the reaction pressure, and separating the oil from the gas by a mercury buffer. The system used is shown in fig 3. The gauge A was connected by steel pressure tubing to a valve B and a pressure vessel C. A Pyrex glass capillary tube D, 1 mm bore, dipped almost to the bottom of this vessel. This capillary

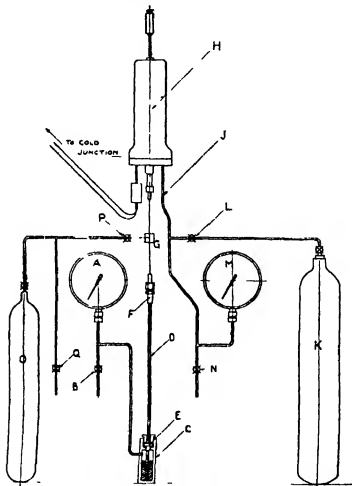


FIG 3

was held by a nut which pressed on a flange E, soldered to the glass. The capillary tube was about a metre long and its upper end was fitted with a soldered-on coupling F, through which it was connected to the reaction bulb, with 0.7 mm bore copper capillary, by way of the T-piece G. The Bourdon tube was evacuated through the valve B, and at the same time the

space over the mercury in the capillary D was evacuated. Refrigerator oil was then allowed to run into the gauge through B, filling up completely all the free space and forcing mercury up the capillary tube to a height approximately barometric. The valve B was then closed. This oil filling made the gauge system so incompressible that a pressure of 80 atmos produced a movement of only 40 cm. in the mercury meniscus in the 1 mm. tube D.

*General Arrangement of the Apparatus*—The complete apparatus is shown in fig. 3. The bomb H was held horizontally on a cradle on a bench about 3 feet high. Passing through its cover plate was a steel pressure tube J, supplying nitrogen. The nitrogen was stored under pressure in the large cylinder K, and the quantity of gas admitted to the bomb was controlled by the fine-adjustment valve L. The pressure of gas in the bomb was shown by the gauge M. A valve N was fitted for blowing off gas or for evacuating the bomb.

The nitrous oxide was stored under pressure in the cylinder O which was connected by steel tubing to a fine-adjustment valve P, leading to the bomb, and a valve Q which was used for blowing off the gas, and for evacuating the reaction bulb. The fine-adjustment valve P controlled the quantity of gas entering the reaction bulb and its speed of entry. It was connected to the T-piece G by 0.7 mm. bore copper capillary tube. This T-piece was connected by similar tubing to the reaction bulb and the glass capillary D. By using this small bore tubing the dead space between the valve P, the reaction bulb, and the mercury meniscus in D was kept below 1 c.c.

The control valves L and P were placed together at shoulder-level on a panel, with the gauges A and M on either side.

Protection against the danger attending a burst of the glass tube D was given by a Triplex glass screen.

*Temperature Measurement and Control*—The Johnson-Matthey platinum-platinum rhodium couple which was used had been calibrated against a platinum resistance thermometer up to about 650° C. and checked at the salt point (801° C.) and gold point (1063° C.). The cold junction was kept at the temperature of melting ice, and the e.m.f. of the couple was measured with a Cambridge Scientific Instrument Company "Workshop potentiometer" which had been altered to give a tapping for a low resistance external galvanometer. The potentiometer dials were set to the required voltage and the current through the furnace was controlled to give no deviation of the galvanometer. A change of one microvolt in the couple e.m.f. produced a deflection of  $1\frac{1}{2}$  mm. By watching for the first signs of a deviation and taking steps to correct it immediately by changing the heating current, it was usually found to be

possible to keep the couple reading constant within less than 2 microvolts, which meant a temperature constant to about  $\pm 0.2^\circ \text{C}$ .

*Experimental Procedure*—After assembling the apparatus the furnace was baked out at the working temperature, and at the same time evacuated through the valve N (fig. 3), by a "Hyvac" pump with a liquid air trap. By doing this a large amount of water and grease was removed from the asbestos lagging, and the danger of shorting the insulated cones by a layer of condensed moisture averted. After this evacuation the bomb was always kept filled with nitrogen.

To prepare for an experiment the furnace temperature was adjusted to the required value and the reaction bulb evacuated through the valves P and Q. Valves P, Q, L, and N were then closed and the valves of the two cylinders N and O were opened. By opening simultaneously the fine adjustment valves P and L, the silica bulb was filled with nitrous oxide at the same time as the bomb was filled with nitrogen and during this filling the pressures of the two gases as shown by the gauges A and M were kept equal. The filling could be carried out at speeds up to 2 atmos per second. When the gas pressure for the experiment was reached the two valves P and L were tightly closed. One observer began immediately to take readings of the gauge A, and of the height of the mercury meniscus in the capillary D, and a second observer started a stop-watch and noted the pressure readings and the time at which they were taken, and also controlled the furnace temperature. A continual light tapping of the gauge was necessary to get the proper reading.

As the reaction proceeded the pressure in the bulb shown by the gauge A rose, and it was necessary to adjust the balancing pressure of nitrogen. This was best done continuously, by adjusting the valve I, to give a steady leakage of gas into the bomb, at a rate corresponding to the rate of increase of pressure in the reaction bulb.

After the reaction had been followed for a sufficient length of time, the gas was blown out by opening valve Q and then slowly releasing valves P and N. When atmospheric pressure was reached, the reading of the gauge A and the height of the mercury column in D was taken. It was then possible to correct the observed readings for the calibration error of the gauge and for the effect of the changing head of mercury in D to obtain the absolute pressures of the gas in the reaction bulb. The results of typical experiments are shown in Table I.

When experiments were made at pressures above approximately 25 atmos there was a tendency for the furnace temperature to fall slightly during the filling, because of the larger quantities of gas which had to be heated. This

fall in temperature was usually less than half a degree, but its effects on the reaction velocity were nullified by starting with the temperature half a degree high.

It is important that the time taken to fill the reaction bulb should be short. If this filling time is an appreciable fraction of the half life, it becomes difficult to decide the initial pressure and starting time of the reaction from the experimental curve, for some of the gas will be reacting and giving a pressure change, although more gas is still entering the bulb. By choosing the working temperature to give half-lives of at least 15 to 20 minutes, the time taken to fill at a speed of 2 atmos per second or faster, becomes such a small fraction of the half-life of the reaction that there is no difficulty in fixing the initial pressure within 0.5%, and the time zero to a second or two.

Good evidence that the time taken by the gas to reach the reaction temperature was very short was provided by the shape of the reaction curves. If the gas had remained below the final reaction temperature for some length of time, the experimental pressure time curve would have shown a pronounced change of curvature near the origin, the points being low, gradually rising to the true reaction curve for a steady temperature as the rising temperature of the gas produced a higher pressure. Actually the experimental curves showed no such effect.

*Source of Nitrous Oxide*—The nitrous oxide was taken from a cylinder of the pure liquid supplied by the British Oxygen Company marked "for inhalation". The gas was found to be 96.7% nitrous oxide, the remainder being nitrogen and oxygen. For many of the experiments in the low pressure apparatus the gas was purified by condensing the nitrous oxide in liquid air and pumping away the permanent gas before evaporating and collecting the pure gas. This purified gas gave the same results as the gas taken directly from the cylinder. This was also the experience of Hinshelwood and Burk, and of Musgrave and Hinshelwood (*loc cit*) who found that even large quantities of air had no effect on the velocity of reaction except at very low pressures of nitrous oxide, e.g., 30 mm Hg.

For the high pressure apparatus cylinder gas was the only convenient source of supply of large quantities of compressed nitrous oxide, and it was used in all the experiments. There was no difficulty in filling the reaction bulb at pressures up to at least 40 atmos, since the vapour pressure from liquid nitrous oxide at room temperatures is above 50 atmos.

*Determination of Nitric Oxide in the Reaction Product*—The amount of nitric oxide formed as a by-product during the high pressure reaction was

measured in a few instances. The method was one of those used by Musgrave and Hinshelwood. When the half-life had been reached the mixture of nitrous oxide and its reaction products was blown into an evacuated, calibrated 3-litre flask, and the temperature and pressure of the gas were measured. In this mixed gas the by-product was present as nitrogen peroxide formed from nitric oxide and oxygen on cooling. It was converted to nitric acid by admitting about 50 c.c. of distilled water to the flask and shaking the contents for half an hour. The acid so formed was titrated with centinormal alkali. Control experiments were made to ensure that no soda was removed from the walls of the flask by the shaking with water, and that the original nitrous oxide shaken with water gave no acid. The percentages of nitric oxide found are given in Table VII, the accuracy is probably about 10%.

*The Temperature and Pressure Ranges Employed*—With the low pressure apparatus experiments were made between 75 mm Hg and 2200 mm Hg at temperatures between 999° and 888° K. The half-lives varied between 120 and 10,000 seconds. The high pressure apparatus was used between  $\frac{1}{2}$  and 40 kg/cm<sup>2</sup> and 840° and 930° K with half-lives varying between 10,000 seconds and about 500 seconds.

Some attempts were made to carry out experiments at pressures as high as 70 kg/cm<sup>2</sup>. To do this it was necessary to warm the supply cylinder to obtain a high enough pressure to fill the reaction bulb, and also to heat the connections to the manometer to prevent the formation of liquid nitrous oxide which would lead to a discrepancy between the observed rate of pressure increase and the true rate of reaction. It becomes necessary to heat the dead space of the system above the critical temperature, 36.5° C., for all experimental pressures much above 40 kg/cm<sup>2</sup>, for when the half-life is reached at about 50–60 kg/cm<sup>2</sup> the pressure is sufficient to liquefy the unreacted nitrous oxide in the dead space if this is at room temperature. These experiments above 40 kg/cm<sup>2</sup> were not successful. On a number of occasions the reaction bulb exploded, probably as a result of contamination of the gas with grease forced from the valve packings by the heating. It is possible that the use of valves packed with fibre or white metal might cure the trouble.

Three of these experiments, although unsatisfactory from the point of view of temperature control and reaction velocity measurement, have provided some of the data for Table VII, which shows the amount of by-product nitric oxide formed during reaction at different pressures.

Table I—Typical Measurements of the Velocity of Decomposition of Nitrous Oxide at High Pressures

857 8° K			887 7° K			875 0° K		
Time from entry of gas	Thermo couple Millivolts	Pressure Kg /cm <sup>2</sup>	Time from entry of gas	Thermo couple Millivolts	Pressure Kg /cm <sup>2</sup>	Time from entry of gas	Thermo couple Millivolts	Pressure Kg /cm <sup>2</sup>
min sec			min sec			min sec		
0 45	5 400	37 52	0 21	5 750	24 62	0 19	5 601	18 41
1 5	5 395	37 57	0 34	5 750	24 77	0 40	5 600	18 46
1 28	5 395	37 62	0 42	5 747	24 78	1 30	5 599	18 66
1 57	5 395	37 72	1 2	5 747	24 88	2 30	5 600	18 70
2 52	5 400	37 93	1 28	5 753	25 05	3 15	5 599	18 85
3 25	5 401	38 03	2 9	5 753	25 25	4 30	5 599	18 98
4 10	5 400	38 18	2 46	5 752	25 47	5 45	5 600	19 09
5 5	5 400	38 33	3 23	5 752	25 73	6 40	5 600	19 25
6 23	5 400	38 53	4 19	5 749	25 96	7 30	5 600	19 35
7 15	5 400	38 68	5 4	5 750	26 21	8 52	5 600	19 51
8 50	5 397	38 94	5 50	5 749	26 42	10 5	5 600	19 66
10 10	5 400	39 10	7 20	5 750	26 83	12 13	5 599	19 82
11 40	5 400	39 44	8 28	5 750	27 14	13 33	5 601	20 07
12 52	5 399	39 59	9 31	5 750	27 30	15 6	5 601	20 22
13 57	5 399	39 64	10 40	5 750	27 56	16 36	5 600	20 36
15 12	5 400	39 94	11 31	5 748	27 82	18 37	5 602	20 61
16 47	5 400	40 20	12 40	5 750	28 07	19 40	5 602	20 71
18 56	5 401	40 50	13 36	5 749	28 21	20 30	5 601	20 81
19 33	5 399	40 65	14 25	5 749	28 41	21 25	5 600	20 90
21 10	5 402	40 84	16 0	5 748	28 75	22 28	5 598	21 00
23 8	5 402	41 14	17 16	5 750	28 99	23 27	5 600	21 10
24 22	5 397	41 29	18 30	5 750	29 22	25 5	5 601	21 23
26 30	5 400	41 54	19 45	5 750	29 41	26 30	5 600	21 36
28 30	5 399	41 84	21 23	5 750	29 71	28 0	5 600	21 60
30 0	5 402	42 03	22 30	5 752	29 99	29 0	5 599	21 60
31 35	5 400	42 23	23 50	5 750	30 09	30 20	5 598	21 70
33 10	5 403	42 43	25 10	5 750	30 28	32 10	5 600	21 86
37 18	5 402	42 93	26 20	5 747	30 43	33 27	5 600	21 95
38 42	5 402	43 13	27 25	5 750	30 58	34 55	5 597	22 00
40 45	5 400	43 32	28 35	5 747	30 75	37 0	5 600	22 24
41 47	5 399	43 43	29 43	5 750	30 87	39 0	5 597	22 33
43 50	5 401	43 53				40 45	5 603	22 48
45 38	5 402	43 78				42 28	5 601	22 57
47 27	5 400	44 03				43 30	5 600	22 66
48 45	5 403	44 23				44 50	5 601	22 76
49 40	5 402	44 33				46 30	5 602	22 85
51 8	5 400	44 43				47 52	5 600	22 94
53 50	5 400	44 73				49 17	5 600	23 08
55 32	5 403	44 94				51 28	5 601	23 18
58 55	5 403	45 14				54 30	5 600	23 27
60 0	5 400	45 39				56 0	5 602	23 36
62 40	5 402	45 39						
64 45	5 399	45 74						
65 52	5 402	45 84						
67 37	5 400	46 00						
69 48	5 400	46 20						
71 45	5 400	46 35						
74 20	5 400	46 55						
75 30	5 400	46 65						
76 20	5 401	46 74						
78 45	5 398	46 93						
81 5	5 399	47 19						

## RESULTS

*The Velocity of Reaction at Pressures up to 2000 mm Hg (2.72 kg/cm<sup>2</sup>) Low Pressure Apparatus*

The results at pressures below 500 mm were obtained with the normal low pressure apparatus, those above this pressure with the low pressure apparatus fitted with closed manometer

Table II

913 5° K		971° K		996 5° K	
Initial pressure mm Hg	Half life, secs	Initial pressure, mm Hg	Half life, secs	Initial pressure, mm Hg	Half life, secs
180	7350	220	910	228	416
400	4760	390	675	375	317
750	3270	832	494.4	829	231.6
1127	2778	988	436.5	932	207.4
1506	2326	1119	399.3	1040	195
1816	2160	1315	360	1126	182.3
2120	1893	1683	307.5	1208	175
		1896	286.2	1278	178.1
		2002	287.5	1385	166.7
				1460	159.7
				1574	148.6
				1728	142.7
				1853	139.2
				1930	131.4
				1983	129.6
				2020	132.2
				2165	122.2

*The Velocity of Reaction at Pressures up to 40 atms High Pressure Apparatus*Table III—Temperature 614.7° C, 887.7° K 1 kg/cm<sup>2</sup> = 0.967 atms

Pressure, kg/cm <sup>2</sup>	Half life, secs	Pressure, kg/cm <sup>2</sup>	Half life, secs
0.36*	13,824	10.63	2,415
0.46*	11,394	12.50	2,093
0.54	9,540	12.84	2,123
2.44	4,815	19.20	1,836
3.40	3,490	24.61	1,542
3.97	3,220	26.67	1,560
4.65	2,970	30.40	1,493
5.64	2,835	32.96	1,482
6.76	2,850	36.45	1,459
8.70	2,425	39.56	1,454

\* Low pressure apparatus



Table IV—Temperature 644 7° C, 917 7° K 1 kg/cm<sup>2</sup> = 0.967 atms

Pressure, kg/cm <sup>2</sup>	Half life, secs	Pressure, kg/cm <sup>2</sup>	Half life, secs
0.51	3,115	7.24	762
2.21	1,680	9.26	711
3.05	1,240	10.78	663
3.59	1,044	12.20	665
4.64	803	12.85	604
5.80	819		

Table V—Temperature 657 5° C, 930 5° K 1 kg/cm<sup>2</sup> = 0.967 atms

Pressure, kg/cm <sup>2</sup>	Half life, secs	Pressure, kg/cm <sup>2</sup>	Half life, secs
0.52	2,195	8.10	534
2.20	957	7.99	473
2.97	787	9.22	429
3.98	658	11.26	398
5.13	585	12.73	348

Table VI—Change of Reaction Velocity with Temperature and Change of Activation Energy with Pressure at Constant N<sub>2</sub>O Concentration

Temperature, °K	Pressure, kg/cm <sup>2</sup>	log <sub>10</sub> (1/t <sub>1</sub> )		Activation energy E, cal/gm mol
		Observed	Calculated	
1 Pressure 37½ kg/cm. <sup>2</sup>				
840.9	37.0	4.0545	4.054	64,900
857.8	37.4	4.383	4.386	
866.2	38.1	4.544	4.548	
875.0	37.85	4.711	4.710	
2 Pressure 18½ kg/cm. <sup>2</sup>				
840.9	18.2	5.948	5.935	63,900
857.8	18.05	4.256	4.261	
866.2	18.6	4.421	4.419	
875.0	18.4	4.5755	4.581	
3 Pressure 0.53 kg/cm. <sup>2</sup> , 390 mm. Hg				
904.3	0.526	4.2645	4.280	61,000
919.6	0.524	4.542	4.528	
934.7	0.536	4.501	4.761	
951.5	0.533	4.796	5.013	
968.5	0.546	5.0075	5.280	
	0.536	5.2640		

Table VI—(continued)

Temperature, °K	Pressure, kg /cm <sup>2</sup>	log <sub>10</sub> (1/t <sub>1</sub> )		Activation energy F cal/gm mol
		Observed	Calculated	
4 Pressure 0.26 kg /cm <sup>2</sup> , 190 mm Hg				
904.3	0.250	4.095	4.109	55,200
934.7	0.256	4.558	4.542	
951.5	0.252	4.7645	4.770	
968.5	0.265	4.770	4.994	
985.8	0.255	4.985	3.213	
	0.258	3.220		
5 Pressure 0.14 kg /cm <sup>2</sup> , 105 mm Hg				
904.3	0.143	5.930	5.943	53,750
	0.137	5.900		
919.0	0.137	4.1845	4.190	
	0.137	4.1725		
951.5	0.143	4.540	4.598	
	0.146	4.631		
968.5	0.141	4.700	4.805	
999.0	0.144	3.192	3.175	
6 Pressure 0.10 kg /cm <sup>2</sup> , 73.5 mm Hg				
916.5	0.10	4.0675	4.050	53,000
934.7	0.10	4.2775	4.298	
951.5	0.10	4.510	4.518	
969.5	0.10	4.750	4.732	
985.8	0.10	4.941	4.941	
999.0	0.10	3.103	3.096	

Table VII—The Amount of By-Product Nitric Oxide formed at various Pressures at 590° C

Initial pressure of N <sub>2</sub> O, kg/cm <sup>2</sup>	11.4	18.2	24.1	33.3	45.3	51.0	63.3
Molecules % NO at the half life	0.11	0.10	0.09	0.07	0.07	0.08	0.09

Nitrous oxide from cylinder (97% N<sub>2</sub>O) 0.0065 mole % NO

## Discussion

The results obtained with the low pressure apparatus, which are shown in fig 4, are clearly in agreement with the observations of Hinshelwood and Burk and Musgrave and Hinshelwood (*loc cit*), since straight-lines can be drawn for  $1/t_1$  against the pressure as far as 2000 mm Hg. From the work of Musgrave and Hinshelwood we know that these lines can be expected to bend sharply towards the origin at some pressure in the neighbourhood of 60 mm. Hg.

Table VIII.—Temperature 888° K

1	2	3	4	5	6	7	8
Pressure kg./cm. <sup>2</sup>	Reacting molecules— 2.5 × colliding molecules	$e - \pi \text{ nr}$	Ratio Col. 2/Col. 3	$f(E)$ $n = 4$	Ratio Col. 2/Col. 5	$f(E)$ $n = 6$	Ratio Col. 2/Col. 7
0.54	1.83 × 10 <sup>-14</sup>	9.3 × 10 <sup>-14</sup>	20.7	144 > 10 <sup>-14</sup>	1.34	11.8 > 10 <sup>-14</sup>	0.164
2.44	0.856 × 10 <sup>-14</sup>	5.01 × 10 <sup>-14</sup>	17.1	79 > 10 <sup>-14</sup>	1.08	6.55 × 10 <sup>-14</sup>	0.120
3.40	0.87 × 10 <sup>-14</sup>	4.17 × 10 <sup>-14</sup>	20.8	66 × 10 <sup>-14</sup>	1.32	5.50 × 10 <sup>-14</sup>	0.158
3.97	0.78 × 10 <sup>-14</sup>	3.80 × 10 <sup>-14</sup>	20.0	62 × 10 <sup>-14</sup>	1.25	5.15 × 10 <sup>-14</sup>	0.152
4.65	0.736 × 10 <sup>-14</sup>	3.39 × 10 <sup>-14</sup>	21.7	54 × 10 <sup>-14</sup>	1.36	4.50 × 10 <sup>-14</sup>	0.164
5.64	0.63 × 10 <sup>-14</sup>	3.16 × 10 <sup>-14</sup>	19.9	50 × 10 <sup>-14</sup>	1.25	4.24 × 10 <sup>-14</sup>	0.148
6.76	0.52 × 10 <sup>-14</sup>	2.86 × 10 <sup>-14</sup>	17.6	47.1 × 10 <sup>-14</sup>	1.10	3.97 × 10 <sup>-14</sup>	0.132
8.70	0.476 × 10 <sup>-14</sup>	2.51 × 10 <sup>-14</sup>	19.0	40.1 × 10 <sup>-14</sup>	1.19	3.40 × 10 <sup>-14</sup>	0.140
10.63	0.39 × 10 <sup>-14</sup>	2.34 × 10 <sup>-14</sup>	16.7	37.6 × 10 <sup>-14</sup>	1.04	3.19 × 10 <sup>-14</sup>	0.122
12.50	0.385 × 10 <sup>-14</sup>	2.04 × 10 <sup>-14</sup>	18.9	32.9 × 10 <sup>-14</sup>	1.17	2.80 × 10 <sup>-14</sup>	0.138
13.84	0.366 × 10 <sup>-14</sup>	2.04 × 10 <sup>-14</sup>	17.9	32.9 × 10 <sup>-14</sup>	1.11	2.80 × 10 <sup>-14</sup>	0.130
19.20	0.404 × 10 <sup>-14</sup>	1.66 × 10 <sup>-14</sup>	24.4	26.9 × 10 <sup>-14</sup>	1.50	2.30 × 10 <sup>-14</sup>	0.176
24.61	0.265 × 10 <sup>-14</sup>	1.45 × 10 <sup>-14</sup>	18.3	24.6 × 10 <sup>-14</sup>	1.08	2.02 × 10 <sup>-14</sup>	0.130
26.67	0.240 × 10 <sup>-14</sup>	1.35 × 10 <sup>-14</sup>	17.8	22.0 × 10 <sup>-14</sup>	1.09	1.90 × 10 <sup>-14</sup>	0.126
30.40	0.223 × 10 <sup>-14</sup>	1.23 × 10 <sup>-14</sup>	18.0	20.0 × 10 <sup>-14</sup>	1.11	1.74 × 10 <sup>-14</sup>	0.128
33.86	0.207 × 10 <sup>-14</sup>	1.18 × 10 <sup>-14</sup>	17.5	19.3 × 10 <sup>-14</sup>	1.07	1.67 × 10 <sup>-14</sup>	0.124
36.45	0.190 × 10 <sup>-14</sup>	1.07 × 10 <sup>-14</sup>	17.7	17.5 × 10 <sup>-14</sup>	1.09	1.62 × 10 <sup>-14</sup>	0.124
39.56	0.176 × 10 <sup>-14</sup>	1.00 × 10 <sup>-14</sup>	17.6	16.4 × 10 <sup>-14</sup>	1.07	1.42 × 10 <sup>-14</sup>	0.124

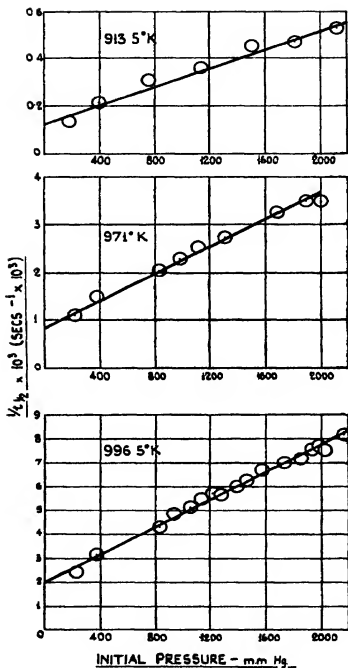


FIG 4

At 2000 mm. Hg, a pressure at which Volmer found an approach to true unimolecular velocity constants, there are no indications whatever of the line  $1/t_1$  against  $p$  tending to become horizontal

The curve for the complete pressure range at 888° K is shown in fig 5, the straight line extending to at least 3 kg/cm.<sup>2</sup> in fig 4 is found to bend over fairly sharply at about 4 kg/cm.<sup>2</sup> to another straight line of smaller slope which persists to about 26 kg/cm.<sup>2</sup> before curving away. Above 30 kg/cm.<sup>2</sup> the half-life is almost independent of pressure

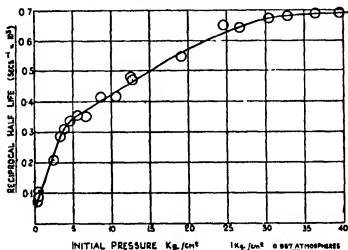


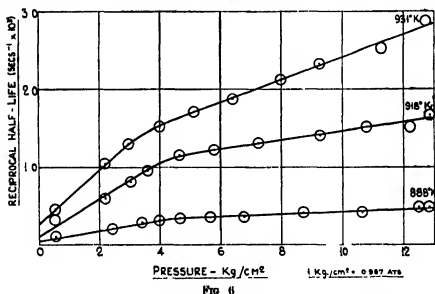
FIG 5

The measurements given in Tables IV and V were made to see whether any shifting of the bend at 4 kg/cm.<sup>2</sup> with change of temperature could be discovered. The curves of fig 6 show that within the range of temperature employed, change of temperature produces no definite change of the pressure at which the bend occurs.

The results of Table VII, which show the amount of nitric oxide formed during reaction, give an assurance that at high pressures catalysis by this by-product is no more serious than it is at low pressures. Musgrave and Hinshelwood found that at an initial pressure of 200 mm Hg of nitrous oxide, about 3½% of nitric oxide was formed during reaction, at 500 mm Hg this had fallen to less than 2%. Table VII shows that the percentage of nitric oxide formed continues to decrease with rising pressure.

The results of measurements made to determine the activation energy are given in Table VI. For each series straight lines are obtained by plotting

$1/K^\circ$  against  $\log 1/t_1$ . The values of  $\log 1/t_1$  calculated back from these straight lines are tabulated in column 4. The inferior accuracy of the two series at the lowest pressures, 75 and 100 mm Hg, is due to the difficulty of measuring accurately with an ordinary manometer, a pressure increase of only 18 to 25 mm Hg. An attempt was made to carry out each series at constant initial concentration of  $N_2O$  rather than at constant initial pressure, by taking into account the variation of pressure with temperature at constant concentration, and adjusting the initial pressures accordingly. The activation energies, calculated on the assumption that the slopes of the lines  $1/K^\circ$  against



$\log 1/t_1$  at constant initial concentration are equal to  $E/R$ , are given in column 5, and are plotted against the pressure in fig 7. In this graph the results of Musgrave and Hinshelwood for zero pressure, Hinshelwood and Burk for pressures between 100 and 400 mm. Hg, and M. A. Hunter for atmospheric pressure are included. Between 0 and 4 kg/cm<sup>2</sup> there is a great increase of activation energy with increasing pressure. Above this pressure there is little further change to 40 kg/cm<sup>2</sup>. These results are at variance with those of Volmer and Nagasako,\* who give an activation energy of 53,000 cal/gm. mol for all pressures up to 10 kg/cm<sup>2</sup>. There is little doubt, however, that fig 7 shows the true behaviour, since it is supported both by the satisfactory results of Table VI and by the results of other workers. It is likely that we

\* 'Z. phys. Chem.', B, vol 9, p 141 (1930)

are concerned with at least two, and possibly more, simultaneous reactions, each with a different activation energy. It remains to be decided whether or not the hypothesis that a unimolecular reaction proceeds alongside a bimolecular one is sufficient. Fig 5 shows that it is not. The straight line for  $1/t - p$  does not continue indefinitely, but bends to another practically straight line of smaller slope, which in turn is succeeded by one which is almost horizontal.

Recent work on the thermal decomposition of acetaldehyde by Hinshelwood and Fletcher\* has produced results which correspond closely to those for

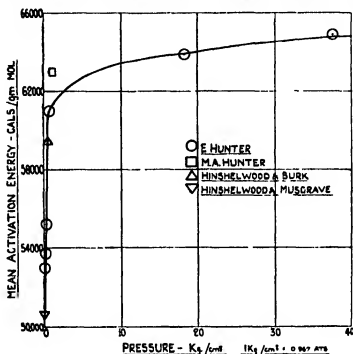


FIG 7

nitrous oxide, except for their much smaller pressure scale. The same type of segmented curve was obtained, with sharp bends at 3, 45, and 250 mm Hg. In order to account for the behaviour of acetaldehyde the theory was put forward that the molecule can be activated in a definite number of different ways, according to the manner in which the energy is divided within the molecule between the different modes of vibration or rotation. In quasi-unimolecular reactions the time lapse between the instant of activation of the molecule by collision and the instant of its decomposition probably arises

\* 'Proc Roy Soc.' A, vol 141, p 41 (1933)

from the necessity for a re-arrangement of the energy within the molecule before a state favouring decomposition is brought about. It is conceivable that the time taken for this re-arrangement will depend on how the energy has been distributed between the modes of vibration according to the particular way in which the molecule has been activated. If a definite number of different ways of activation are possible they will lead to the same definite number of independent quasi-unimolecular reactions.

In their paper, Hinshelwood and Fletcher made a rough analysis of a curve of the type of fig 5 and succeeded in showing that such a curve represents the combined effect of three separate quasi-unimolecular reactions.

For each reaction an equation of the type

$$K_1 p^2 - K_2 p a - K_3 a = 0,$$

holds, where  $K_1 p^2$  is the rate of activation by collision,  $K_2 p a$  the rate of deactivation of active molecules by collision with normal molecules, and  $K_3 a$  the rate of decomposition of activated molecules, or rate of reaction. If there are three types of activated state the observed velocity is

$$-dp/dt = \frac{K_1 p^2}{1 + K_2 p/K_3} + \frac{K'_1 p^2}{1 + K'_2 p/K'_3} + \frac{K''_1 p^2}{1 + K''_2 p/K''_3}$$

If the three types of activated molecule have fairly widely different probabilities of reaction,  $K_3 \gg K'_3 \gg K''_3$ , and it can be shown that the curve for  $1/t_1$  against the initial pressure starts from the origin as

$$1/t_1 = (K_1 + K'_1 + K''_1) p_0$$

This is the line OA of fig 8. It is succeeded by a practically straight line of slope approximately  $(K_1 + K'_1)$  which makes an intercept of about  $K''_1 K''_3 / K''_2 \log 2$ , on the axis of  $1/t_1$ . This line, AB in fig 8, represents the reaction velocities at pressures between 50 and about 3000 mm Hg, and its intercept is that which appears in figs 4 and 5. In this region one of the three reactions has reached its limiting velocity constant. At a higher pressure the second reaction reaches its limiting velocity constant, and the line of slope  $(K_1 + K'_1)$  is followed by a line of slope  $K_1$ , directed to a still larger intercept on the axis of  $1/t_1$ . This third line is BC in fig 8 which eventually becomes practically horizontal as the rising pressure causes the third reaction to approach its limiting velocity constant.

The methods of this analysis give for the nitrous oxide decomposition the three quasi-unimolecular curves OEF, OGH, and OJK of fig 8. Although it



is clear that three separate curves can account for the shape of the experimental curve OABCD, fig 8 does not show more than the approximate magnitudes and shapes of the three curves, since the limiting velocity constants  $L_F$ ,  $L_H$ , and  $L_K$  are not sufficiently widely separated to fulfil the condition  $K_1 > K'_2 > K''_3$ , which is necessary before the above analysis can be expected to hold almost quantitatively

It is now possible to explain the activation energy curve of fig 7. Each of the three reactions has its own activation energy. At low pressures ( $< 50$  mm

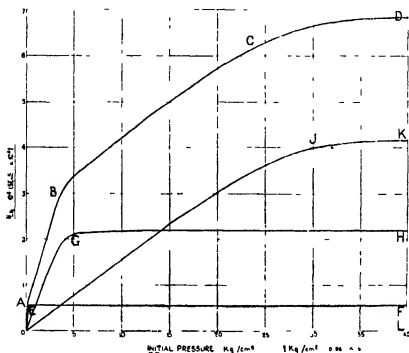


FIG 8

Hg) the low pressure reaction OEF of fig 8 contributes a greater fraction to the decomposition than either the medium pressure reaction OGH, or the high pressure reaction OJK, but as the pressure rises these latter two reactions participate to a greater and greater extent. The curve for the average activation energy consequently shows a rapid change in  $E$  with increasing pressure. At about 5 kg/cm<sup>2</sup>, however, both the low pressure and the medium pressure reactions are giving their maximum possible reaction velocities, and their combined contributions are quite a large fraction of the total. The increase in

velocity on increasing the pressure above about 5 kg/cm.<sup>2</sup> is due solely to the growth of the high pressure reaction OJK of fig 8 and the measured mean activation energy no longer shows such a rapid change with increasing pressure. Above a pressure of about 30 kg/cm.<sup>2</sup> the limiting velocity constants of all three reactions are reached and the observed activation energy becomes independent of pressure, since the relative contributions of the three reactions do not vary with pressure.

The absolute values of all three activation energies cannot be deduced from fig 7 since each may itself vary continuously with pressure. It is apparent that the low pressure reaction is associated with an activation energy of about 50,500 cal/gm. mol. The activation energy of the high pressure reaction must be above 65,000 cal/gm. mol., and the value for the medium pressure reaction is probably between 50,500 and 62,000 cal/gm. mol. Since the first reaction with activation energy 50,500 cal/gm. mol. gives a maximum velocity less than that for the second reaction with activation energy approximately 60,000 cal/gm. mol., and this second reaction in its turn gives a maximum velocity less than that of the third reaction with an activation energy greater than 65,000 cal/gm. mol., the probability of the decomposition of the activated molecule of the first reaction is smaller than that of the second, which again is smaller than that of the third reaction with the highest of the three activation energies.

In Table VIII the results of some calculations based on the data of Tables III and VI are given. Since the reaction is a compound one it was not justifiable to assume that the number of molecules reacting per cubic centimetre per second at any given pressure could be calculated directly from the particular value of the unimolecular velocity "constant"  $1/n_2/t_1$ , or the bimolecular constant,  $1/at_1$ , at that pressure. However, the tangent  $dp/dt$  at the time origin of each reaction curve can be used to calculate the number of molecules reacting, since the pressure increase is brought about by the formation of three molecules in place of two of nitrous oxide. It was found that the number calculated in this way always agreed to within 10% of the number calculated from the different values of  $1/n_2/t_1$  (the unimolecular "constant") over the pressure range. The numbers calculated in the latter way were used in Table VIII since the values of  $t_1$  were known more accurately than the values of the tangents at the very start of the reactions, which were not easy to determine accurately.

In column 2 of Table VIII is shown the ratio, reacting molecules/(2.5 × calc. no. of colliding molecules). The number of collisions was calculated

on the assumption that the diameter of the nitrous oxide molecule is  $3.32 \times 10^{-8}$  cm. The factor 2.5 is used to compensate for the increased number of collisions due to the higher than average velocity of the high translational energy molecules which are likely to be concerned in activating collisions and for the possibility that a particular orientation of the molecules at the moment of impact is necessary.

The values calculated for the expression  $e^{-E/RT}$  from the data of Table VI are given in column 3 of Table VIII, and it will be seen that activation energy associated with only two square terms can only account for one-twentieth of the observed reaction, as is shown in column 4.

The fraction of the molecules possessing energy greater than  $E$ , distributed between a number of degrees of freedom is given as a sufficient approximation by the expression

$$\frac{e^{-\left(\frac{E + (\frac{1}{2}n - 1)RT}{RT}\right)}}{\frac{1}{2}n - 1} \left[ \frac{E/RT + (\frac{1}{2}n + 1)}{\frac{1}{2}n - 1} \right]^{1/2n-1}$$

where  $n$  represents the number of "square terms" between which the energy is distributed. This expression worked out for nitrous oxide, gives for  $n = 4$  and  $n = 6$ , the values which appear in columns 5 and 7, respectively of Table VIII. The ratios in columns 6 and 8 show that the rate of activation can be equated to rate of reaction if  $n = 4$  and is greater than the rate of reaction if  $n = 6$ . This implies a distribution of energy between two, or at most three degrees of freedom, so that activation energy can be distributed between translational energy and one or perhaps two modes of vibrational energy within the molecule. It is now fairly generally accepted that the atoms of the nitrous oxide molecule lie in a straight line in the order NNO, two or even three degrees of freedom is therefore not an unreasonable number.

Since the observed reaction can be regarded as the resultant of three quasi-unimolecular reactions, the thermal decomposition of nitrous oxide is principally concerned with the isolated molecule, which presumably breaks up according to the scheme  $N_2O \longrightarrow N_2 + O$ . This free oxygen atom must react rapidly, either with a second free atom,  $O + O \longrightarrow O_2$ , or with a fresh nitrous oxide molecule  $O + N_2O \longrightarrow N_2 + O_2$ , and the measured pressure increase will be that corresponding to the stoichiometric equation  $2N_2O \longrightarrow 2N_2 + O_2$ .

The author wishes to thank Mr C. N. Hinshelwood for his help and advice throughout the work, and Mr W. R. D. Manning, of the Department, for his help in the design of the apparatus.

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### *Summary*

Apparatus for the study of gas reactions by the static method at pressures up to 100 atms and at temperatures up to 900° C is described

Results of measurements of the velocity of thermal decomposition of nitrous oxide at pressures up to 40 atms are given. In conjunction with the work of Musgrave and Hinshelwood\* they show clearly that the observed reaction is the resultant of three separate quasi-unimolecular reactions which become independent of pressure at approximately 0.08, 5, and 30 atms, respectively.

The average activation energy of the reaction has been measured at different pressures and is found to rise rapidly from the value 50,500 cal/gm. mol at zero pressure until a pressure of about 6 atms is reached. As the pressure rises above 6 atms it becomes less and less dependent upon pressure, and is practically independent of pressure above 30 atms.

The behaviour of the reaction suggests that there are three separate modes of activation, each with a different mean activation energy and a different probability of decomposition of the activated molecule.

\* 'Proc. Roy. Soc.,' A, vol. 135, p. 23 (1932)

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## *A Relativistic Basis of the Quantum Theory*

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### *Introduction*

In a previous communication\* it was pointed out that the problem of the quantum equations could be regarded as having developed into a search for a modification of the Riemannian covariant differentiation.

This view of the problem appears from many of the more recent publications, some of which are quoted below †.

Various underlying geometrical or analytical theories have been employed to determine the modification of the Christoffel bracket expression, which we write here  $\Gamma_{\mu\nu}^\sigma$ , but in general the effect is to introduce a new quantity  $T_{\mu\nu}^\sigma$  which must be added to  $\Gamma_{\mu\nu}^\sigma$ . The construction of  $T_{\mu\nu}^\sigma$  depends upon the theory adopted, and it is sometimes symmetrical in  $\mu, \nu$  and sometimes not.

Frequently the attempts have been successful to the extent of uniting gravitational and electromagnetic theory but the quantum equations, though fitting in well with the particular geometrical background, stand aside somewhat and are not absorbed completely into the scheme. As an example of this Fisher and Flint (*loc. cit. infra*) found that Dirac's first order equations could be expressed in the form of a vanishing divergence and from the equations so obtained the second order wave equation containing the "spin" terms was deduced without the introduction of further hypothesis. But this method merely placed the equations into the scheme and it was only by a somewhat indirect process that they could be shown to have a geometrical significance ‡.

Nevertheless, the actual form of the first order equations in this notation raises the hope of a possible union of gravitational, electromagnetic and quantum theories. The present paper is a result of this discovery. The underlying idea of this earlier attack on the problem was simple. The track of a charged particle in a gravitational and electromagnetic field was regarded as a null geodesic. Associated with this track a set of equations was adopted,

\* Flint, 'Proc. Roy. Soc.' A, vol. 141, p. 370 (1933).

† Flint, 'Proc. Roy. Soc.' A, vol. 117, p. 630 (1928), Fisher and Flint, 'Proc. Roy. Soc.' A, vol. 126, p. 644 (1930), Fock, 'Z. Physik,' vol. 57, p. 261 (1929), Schroedinger 'Sitzber. Preuss. Akad. Wiss.' No. 11, p. 106 (1932), Pauli, 'Ann. Physik,' vol. 18, pp. 305, 338 (1933), Schouten and van Dantzig, 'Z. Physik,' vol. 78, p. 639 (1932).

‡ Fahmy, 'Proc. Phys. Soc.', vol. 44, p. 368 (1932).

based on the analogy that exists between the track of a photon and Maxwell's equations. The equations so obtained turned out to be in the form of the quantum equations.

In the light of the present study of the problem we shall have to regard this as an approximate theory, but it may be that in this close analogy to the wave theory of light we have the basis of the wave theory of matter. We may see in examining the extent to which the analogy may be extended just how far we can go with the wave theory of matter.

The other view of the quantum problem, which we suggested\* was that it has become a search for quantities more fundamental in character than the  $g_{\mu\nu}$  in the expression,  $ds^2 = g_{\mu\nu} dx^\mu dx^\nu$ , for the line element.

The first to express this definitely appears to have been Tetrode†. Fock (*loc cit*) worked out a detailed theory, using a relation of Tetrode's shortly afterwards.

More recently Schroedinger (*loc cit*) has developed a theory of invariance with a generalized set of quantum equations, with which we shall have occasion to compare the results of this paper.

The basis of this line of attack is the introduction of matrices  $\alpha_m$ , which are more general than those originally introduced by Dirac and which satisfy the equations

$$\alpha_m \alpha_n + \alpha_n \alpha_m = g_{mn} I, \quad (1)$$

where  $I$  is the unit matrix.

This method has much in common in its form with Einstein's theory of parallelism,‡ which we showed lent itself readily to the expression of the quantum equations as a vanishing divergence§. It was therefore thought that the geometrical representation might succeed best in the framework of that theory and a study of the question was published||. A close analogy was discovered with Schroedinger's method but nothing emerged from the theory which would help to decide upon the particular type of space-time geometry appropriate to natural phenomena. The mathematical form of the theory appears to permit of some variation in this particular.

We must just mention here the work of Veblen and Hoffmann and the very general theory of Schouten and van Dantzig. The latter have given a very

\* 'Proc Roy Soc,' A, vol. 141, p. 370 (1933)

† 'Z. Physik,' vol. 50, p. 336 (1928)

‡ Einstein, 'SitzBer. Preuss. Akad. Wiss.,' No. 17, p. 217 (1928)

§ Flint, 'Proc Roy Soc,' A, vol. 121, p. 676 (1928)

|| *Ibid.*, vol. 141, p. 363 (1933)

general formulation of the field theory, and it appears to include most of the important notations already proposed

### *The Basis of the Present Theory*

The method we adopt here is based on a modification of the Riemannian covariant derivative

Instead of taking this to be

$$\frac{\partial A_\mu}{\partial x^\nu} - \Gamma_{\mu\nu}^\alpha A_\alpha, \quad (2)$$

we shall suppose it to be

$$\frac{\partial A_\mu}{\partial x^\nu} - \Delta_{\mu\nu}^\alpha A_\alpha, \quad (3)$$

and we shall write

$$\Delta_{\mu\nu}^\alpha = \Gamma_{\mu\nu}^\alpha + T_{\mu\nu}^\alpha \quad (4)$$

$T_{\mu\nu}^\alpha$  will not be supposed symmetrical in  $\mu, \nu$  as  $\Gamma_{\mu\nu}^\alpha$  is, but we shall divide it into an antisymmetrical part  $\Lambda_{\mu\nu}^\alpha$  and a symmetrical  $\Theta_{\mu\nu}^\alpha$ , writing

$$T_{\mu\nu}^\alpha = \Lambda_{\mu\nu}^\alpha + \Theta_{\mu\nu}^\alpha \quad (5)$$

It will be seen that we are using the same notation as Einstein employed. This is for the reason mentioned above that his theory of parallelism suggests the mathematical form appropriate to our purpose. We do not adopt the system of parallelism introduced by him. We shall, at any rate, have no occasion to refer to it. The idea is to leave open questions of this sort in order to free the notation of the limitations which geometry imposes on the form of  $T_{\mu\nu}^\alpha$ . It will be noticed that we have left ourselves unrestricted with regard to symmetry.

It must also be made clear that we propose to make use of a system of five co-ordinates,  $x_\mu$ , ( $\mu = 1, 2, 3, 4, 5$ ). Every particle in our theory has five degrees of freedom. This is a point which has not met with general acceptance on the ground that the space-time of Physics is four dimensional and no definite meaning can be given to the fifth co-ordinate.

Our use of the fifth co-ordinate here is solely to enable us to make an appeal to mathematical form. If we are correct in the sense that our results correspond accurately with those of experiment, we can never presume to say more than that the physical world can be described as if it were five dimensional. The success of the four-dimensional theory of the universe allows us to say no more than this with regard to the four-dimensional world. One may only exclaim in the excitement of discovery that space and time have henceforth vanished to shadows.

But as we have stated above, it is not our intention to insist upon the geometrical aspect of the theory

The more serious objections to the theory in the form of Kahira and Klein are that the general covariance is destroyed by the "cylindrical" condition that the  $q_{\mu\nu}$  do not contain  $x^5$  and that  $q_{55}$  is taken as constant

For this reason many writers have sought to maintain a four-dimensional continuum, introducing five variables as auxiliary quantities, related to the four co-ordinates, e.g., Pauli's method of homogeneous co-ordinates makes the space-time co-ordinates homogeneous functions of zero degree in the homogeneous variables

In the theory of Eddington and Mayer\* vectors are introduced which are related to four-dimensional vectors, but the co-ordinate system is four-dimensional. This theory is a unitary theory of gravitation and electricity

Such theories necessarily include many definitions and assumptions about the quantities introduced and extensions to the inclusion of the quantum theory involve some complications

We shall try to meet these objections by developing a more general theory than has been attempted hitherto. It will be seen that the results obtained follow very naturally from the application of the methods of the theory of relativity

We shall then regard the special use of the cylindrical condition and the assumption about the way  $x^5$  occurs in the functions as an approximation required by our need to eliminate  $x^5$ , in interpreting our results in the light of our present knowledge of physical phenomena. We have to adopt a somewhat similar process in writing  $x^4 = ut$ , when we use the four co-ordinates  $x^1, x^2, x^3, x^4$ , in the theory of relativity. We cannot, of course, claim that  $x^5$  is in such a good case in this respect as  $x^4$ . This is our apology, if apology be needed, for the practical use of  $x^5$  in this restricted way

### *Curvature*

Following the suggestion that the geometrical notation gives, we shall describe the quantity  $P_{\mu\nu\sigma}^{\phantom{\mu\nu\sigma}\rho}$  as the curvature tensor. In the notation of (3) the expression for this quantity is

$$P_{\mu\nu\sigma}^{\phantom{\mu\nu\sigma}\rho} = \Delta_{\mu\sigma}^{\phantom{\mu\sigma}\nu\rho} - \Delta_{\mu\nu}^{\phantom{\mu\nu}\sigma\rho} + \Delta_{\rho\sigma}^{\phantom{\rho\sigma}\mu\nu} - \Delta_{\rho\nu}^{\phantom{\rho\nu}\mu\sigma} \quad (6)$$

The use of the comma before a suffix denotes ordinary differentiation with

\* 'SitzBer Preuss Akad. Wiss.', No 25, p 541 (1931)



respect to the corresponding variable, e.g.,  $\Delta_{\mu\sigma}{}^\nu = \frac{\partial \Delta_{\mu\sigma}}{\partial x^\nu}$ . We shall denote

covariant differentiation by the symbol  $\nabla$ . It must be understood that the Greek affixes can have the values 1-5.

If the substitution of (4) be made it follows that

$$P_{\mu,\sigma}{}^\nu = R_{\mu,\sigma}{}^\nu - T_{\mu,\sigma}{}^\nu + T_{\mu\sigma}{}^\nu - T_{\rho\sigma}{}^\nu T_{\mu}{}^\rho + T_{\rho\nu}{}^\sigma T_{\mu}{}^\rho \quad (7)$$

This may be compared with the corresponding expression in the theory of relativity\* where the quantity  $T_{\mu,\sigma}{}^\nu$  is symmetrical in  $\mu, \nu$ . The contracted curvature tensor obtained by writing  $\varepsilon = \sigma$  is

$$P_{\mu\nu} = R_{\mu\nu} - T_{\mu,\sigma}{}^\sigma + T_{\mu\sigma}{}^\sigma - T_{\rho\sigma}{}^\sigma T_{\mu}{}^\rho + T_{\rho\nu}{}^\sigma T_{\mu}{}^\rho \quad (8)$$

### The Value of $T_{\mu,\sigma}{}^\nu$ and the Second Order Equation

In imposing a restriction upon this quantity we shall be guided again by the geometrical methods.

One suggestion that arises is to write

$$T_{\mu\nu}{}^\sigma = \gamma_{\mu\nu} \Pi^\sigma - \gamma_\mu{}^\sigma \Pi_\nu - \gamma_\nu{}^\sigma \Pi_\mu \quad (9)$$

where the quantities  $\gamma_{\mu\nu}$ ,  $\gamma_\mu{}^\sigma$ ,  $\gamma_\nu{}^\sigma$  have their usual meanings. We write for the five-dimensional line element

$$ds^2 = g_{\mu\nu} dx^\mu dx^\nu, \quad (10)$$

so that the  $\gamma_{\mu\nu}$  are analogous to the  $g_{\mu\nu}$  of Riemannian geometry.

This suggestion comes directly from the theory of Eddington and Weyl\*. It is the suggestion made formerly† which proved very fruitful in the expression of Dirac's equations as a vanishing divergence. But we find it too restricted for our present purpose. It may be noted that in this form  $T_{\mu\nu}{}^\sigma$  is symmetrical in  $\mu, \nu$ .

We return rather to a form used in Einstein's theory‡

$$T_{\mu\nu}{}^\sigma = \Lambda_{\mu\nu}{}^\sigma - \gamma_\mu{}^\sigma \gamma_{\nu\rho} \Lambda_{\tau}{}^\rho - \gamma_\nu{}^\sigma \gamma_{\mu\rho} \Lambda_{\tau}{}^\rho \quad (11)$$

$\Lambda_{\mu\nu}{}^\sigma$  is antisymmetrical in  $\mu, \nu$  and it will be seen that the remaining terms on the right, taken together, are symmetric in  $\mu, \nu$ . Thus the value of  $\Theta_{\mu\nu}{}^\sigma$  is the sum of these terms.

\* Eddington, "Mathematical Theory of Relativity," 2nd ed., p. 218.

† Fisher and Flint, *loc. cit.* (1930).

‡ See also Flint, "Proc. Roy. Soc., A," vol. 141, p. 371 (1933).

We have thus assumed that  $T_{\mu\nu}$  depends upon a single quantity with components  $\Lambda_{\mu\nu}$  except, of course, for the  $\gamma_{\mu\nu}$ .

We can without very much calculation substitute for  $T_{\mu\nu}$  in the contracted curvature (8). The work is rendered much lighter if we remember that the covariant derivatives of the  $\gamma_{\mu\nu}$  vanish.

We find that it is important for us to consider the quantity  $\gamma^{\mu\nu}P_{\mu\nu}$  so that we can omit the separate calculation of  $P_{\mu\nu}$ .

We find that

$$\gamma^{\mu\nu}P_{\mu\nu} = \gamma^{\mu\nu}R_{\mu\nu} - 4\gamma^{\mu\nu}\Lambda_{\sigma\mu}{}^{\sigma}{}_{\nu} + 4\gamma^{\sigma\tau}\Lambda_{\sigma\sigma}{}^{\tau}\Lambda_{\tau\tau}{}^{\mu}{}_{\mu} + \gamma^{\sigma\tau}(\gamma^{\mu\nu}\gamma_{\mu\nu} + 2\delta_{\mu}{}^{\mu}\delta_{\nu}{}^{\nu})\Lambda_{\tau\sigma}{}^{\mu}\Lambda_{\mu\tau}{}^{\nu} \quad (12)$$

Let us write  $\Lambda_{\sigma\mu}{}^{\sigma}{}_{\nu} = \frac{\alpha}{\psi} \frac{\partial\psi}{\partial x^{\mu}}$

Then

$$\begin{aligned} \gamma^{\mu\nu}\Lambda_{\sigma\mu}{}^{\sigma}{}_{\nu} &= \alpha\gamma^{\mu\nu}\left\{\frac{\partial}{\partial x^{\sigma}}\left(\frac{1}{\psi}\frac{\partial\psi}{\partial x^{\mu}}\right) - \Gamma_{\mu\nu}{}^{\sigma}\left(\frac{1}{\psi}\frac{\partial\psi}{\partial x^{\sigma}}\right)\right\} \\ &= \frac{\alpha}{\sqrt{\gamma}}\frac{\partial}{\partial x^{\sigma}}\left\{\sqrt{\gamma}\gamma^{\mu\nu}\frac{1}{\psi}\frac{\partial\psi}{\partial x^{\mu}}\right\} \\ &= \frac{\alpha}{\psi}\frac{1}{\sqrt{\gamma}}\frac{\partial}{\partial x^{\sigma}}\left\{\sqrt{\gamma}\gamma^{\mu\nu}\frac{\partial\psi}{\partial x^{\mu}}\right\} - \frac{\alpha}{\psi^2}\gamma^{\mu\nu}\frac{\partial\psi}{\partial x^{\mu}}\frac{\partial\psi}{\partial x^{\sigma}} \\ &\quad - \frac{\alpha}{\psi}\frac{1}{\sqrt{\gamma}}\frac{\partial}{\partial x^{\sigma}}\left\{\sqrt{\gamma}\gamma^{\mu\nu}\frac{\partial\psi}{\partial x^{\mu}}\right\} - \frac{\gamma^{\mu\nu}}{\alpha}\Lambda_{\sigma\mu}{}^{\sigma}{}_{\nu}\Lambda_{\tau\sigma}{}^{\mu}\Lambda_{\mu\tau}{}^{\nu} \end{aligned}$$

If  $\alpha$  be made equal to  $-1$ , we eliminate the third term on the right of (12), which may now be written

$$\frac{1}{\psi}\bigcirc\psi + \frac{1}{4}\gamma^{\sigma\tau}(\gamma^{\mu\nu}\gamma_{\mu\nu} + 2\delta_{\mu}{}^{\mu}\delta_{\nu}{}^{\nu})\Lambda_{\tau\sigma}{}^{\mu}\Lambda_{\mu\tau}{}^{\nu} + \frac{1}{4}\gamma^{\mu\nu}R_{\mu\nu} = \frac{1}{4}\gamma^{\mu\nu}P_{\mu\nu} \quad (13)$$

$\gamma^{\mu\nu}R_{\mu\nu}$  is the Gaussian curvature, i.e., if we regard space as five-dimensional Riemannian space, we can speak of it as the curvature of that space. We shall denote it by  $R$  and it corresponds to the  $G$  of relativity theory. We shall denote  $\gamma^{\mu\nu}P_{\mu\nu}$  by  $P$ .

The symbol  $\bigcirc\psi$  is familiar in the theory of Klein. It corresponds to the Laplacian operator and in Klein's theory

$$\bigcirc\psi = 0 \quad (14)$$

is Schrodinger's equation for an electron without spin in a gravitational and electromagnetic field.

We make the suggestion that the complete second order wave equation of the quantum theory in its relativistic form is (13)

The reason for this suggestion will be made clear by considering the approximate form obtained by the usual method of eliminating  $x^5$

### Reduction of the General Equation

In order to compare (13) with the equations proposed hitherto we shall adopt the Klein-Kaluza values of the  $\gamma_{\mu\nu}$

If  $m, n$ , represent values 1-4, the values are

$$\left. \begin{aligned} \gamma_{mn} &= g_{mn} + \gamma_{55}\alpha^2\phi_m\phi_n, & \gamma_{m5} &= \gamma_{5m} = \gamma_{55}\alpha\phi_m, \\ \gamma^{mn} &= g^{mn}, & \gamma^{m5} &= \gamma^{5m} = -\alpha\phi^m, & \gamma^{55} &= \alpha^2\phi_m\phi^m + \frac{1}{\gamma_{55}}, \\ \gamma_{55}\alpha^2 &= -\frac{16\pi K}{c^2}, \end{aligned} \right\} \quad (15)$$

$\phi_m$  is a component of the electromagnetic potential and  $K$  is the Newtonian constant of gravitation

With this set of values the track of a charged particle is a geodesic in five-dimensional space and by writing  $\alpha = e/mc$ , it becomes a null geodesic \*. But however suggestive this may be it appears to be numerically incorrect in further applications of the theory

We shall further make use of de Broglie's suggestion that  $x^5$  occurs in such a way in those quantities that depend upon it that differentiation with respect to this variable is equivalent to multiplication  $2\pi i e/h\alpha$ ,  $\alpha$  being the constant in (15) and  $e$  the value of the fundamental electric charge

If  $\alpha = e/mc$ , this is equivalent to multiplication by  $2\pi i mc/h$ , which is often taken to replace  $\partial/\partial x^5$ , but this makes the theory too restricted and de Broglie has considerably helped the theory by not introducing the mass in this operation

With these limitations and taking the value of  $\sqrt{-\gamma}$  as unity, as in the theory of relativity, we obtain

$$\square\psi = \square^2\psi - \frac{4\pi e}{h} \frac{\partial\psi}{\partial x^m} - \frac{4\pi^2}{h^2} \left( \frac{e^2}{c^2} \phi_m\phi^m + \frac{e^2}{\alpha^2\gamma_{55}} \right) \quad (16)$$

We notice that additional terms occur in (13), one of these must represent the

\* Fisher, 'Proc Roy Soc,' A, vol 123, p 489 (1929)

spin term and another the mass of the particle considered. The third term  $\gamma^{\mu\nu} R_{\mu\nu}$  is the curvature, and its value is known\*

$$\gamma^{\mu\nu} R_{\mu\nu} = G - \frac{8\pi K}{c^2} H \quad (17)$$

$$H = F^{\mu\nu} F_{\mu\nu} \quad (18)$$

$$F_{\mu\nu} = \frac{\partial \phi_\nu}{\partial x^\mu} - \frac{\partial \phi_\mu}{\partial x^\nu}, \quad (19)$$

$G$  is the four-dimensional curvature

The occurrence of the curvature in the wave equation has been previously noticed by Schroedinger (loc cit) and a comparison of (13) with his equation (74) reveals a striking resemblance

An interesting point in connection with our equation is that we have not yet introduced the mass. This mass term has caused a difficulty in the derivation of a general equation, for it has always been necessary to introduce the mass of the electron or proton arbitrarily

There seems no doubt here how it must be introduced. In the notation of Klein the mass of a particle takes a secondary position and is replaced by a quantity  $I/c$ ,

$$I^2 = m^2 c^2 + \frac{e^2}{\alpha^2 \gamma_{55}} \quad (20)$$

In the case when the geodesic is a null geodesic and  $\gamma_{55} = -1$ ,  $I$  is zero. Thus we have the further analogy between mass point and photon. If for the one is zero, the rest mass is zero for the other. But we cannot accept this as a general result. We must admit a non-vanishing value for  $I$ . We shall suppose that

$$i\gamma^{\mu\nu} P_{\mu\nu} = -\frac{4\pi^2 I^2}{\hbar^2} = -\frac{4\pi^2}{\hbar^2} \left( m^2 c^2 + \frac{e^2}{\alpha^2 \gamma_{55}} \right) \quad (21)$$

Thus the quantity, which we might call the generalized curvature, is determined by the mass and charge of the particle situated at the point where the curvature is measured

This geometrical analogy gives a satisfactory way of introducing the mass into our equation

The negative sign appears to be rather arbitrarily chosen but the deciding factor is that we know from Schroedinger's equation that  $4\pi^2 m^2 c^2 / \hbar^2$  occurs with a positive sign when taken over to the side of  $\square \psi$ . It will be seen at

\* Rosenfeld, 'Bull. Acad. Roy. Belgique,' vol. 13, p. 304 (1927)

the same time that  $e^2/\alpha^2\gamma_{ss}$  cancels from the equation taking with it the gravitational constant. Gravitation is represented in this part of the equation by the curvature  $G$ . We have shown previously that the "spin" terms occur in the form of a term

$$-\frac{2\pi e}{h}(H_1A^{23}+H_2A^{31}+H_3A^{12}-iE_1A^{14}-iE_2A^{24}-iE_3A^{34}),$$

and in Schroedinger's equation (74) of his paper (*loc. cit.*) they are represented by  $-\frac{1}{2}f_{kl}\gamma^{kl}$ .

Our additional term is of this type and we shall write it in the form

$$-\frac{2\pi e}{h}F_{\mu\nu}A^{\mu\nu}$$

to bring it into the same notation.

Thus the reduced equation becomes

$$\frac{1}{\psi}\square^2\psi-\frac{4\pi e}{h}\left(\phi^m\frac{1}{\psi}\frac{\partial\psi}{\partial x^m}-\frac{2\pi e}{h}F_{\mu\nu}A^{\mu\nu}+\frac{1}{2}\left(G-\frac{4\pi K}{c^2}H\right)+\frac{4\pi^2}{h^2}\left(m^2c^2-\frac{e^2}{c^2}\phi_m\phi^m\right)\right)=0 \quad (22)$$

We can write this in a rather more general form by giving to the  $\square^2\psi$  of (16) the value it has in (13). We then obtain

$$\frac{\gamma^{\mu\nu}}{\psi}\left(\frac{\partial^2\psi}{\partial x^\mu\partial x^\nu}-\gamma^{\mu\nu}\frac{\partial\psi}{\partial x^\alpha}\right)-\frac{2\pi e}{h}F_{\mu\nu}A^{\mu\nu}+\frac{1}{2}\left(G-\frac{4\pi K}{c^2}H\right)+\frac{4\pi^2}{h^2}\left(m^2c^2-\frac{e^2}{16\pi K}\right)=0 \quad (23)$$

This form should be compared with an equation of the electron without spin given by de Broglie\*.

The term in the curvature is of very great theoretical interest but much too small, compared with the term in  $m$ , to effect an appreciable result.

The importance of the derivation of (13) or (23) is that it arises quite naturally from relativistic methods and is nothing else than the equation of curvature

### The Relation to Matrix Theory

So far no matrices have been used in the theory. The equation (23) contains a scalar quantity  $\psi$  introduced through  $\Lambda_{\alpha\mu}\psi$  and is of the same form as the earlier Schroedinger equations.

\* 'J. Phys. Rad.', vol. 3, p. 65 (1937).

Matrices are introduced in attempting to replace the  $\gamma_{\mu\nu}$  by more fundamental quantities

The way this is done is indicated in equation (1)

As we have previously pointed out there is a strong resemblance between these quantities and Einsteins'  $h_{\mu\nu}$  of the theory of parallelism. We therefore subject the  $\alpha_\mu$  to the following condition

$$\frac{\partial \alpha_\mu}{\partial x^\lambda} - \Gamma_{\mu\lambda}^\sigma \alpha_\sigma - T_{\mu\lambda}^\sigma \alpha_\sigma = 0 \quad (24)$$

This may be compared with equation (14) of an earlier paper \*

Bargmann† has pointed out that this leads directly to Schroedinger's method, if it be postulated that  $T_{\mu\lambda}^\sigma \alpha_\sigma$  is of such a form that

$$T_{\mu\lambda}^\sigma \alpha_\sigma = \Gamma_\lambda^\sigma \alpha_\mu - \alpha_\mu \Gamma_\lambda^\sigma \quad (25)$$

Equation (24) is similar to the Riemannian relation

$$\gamma_{\mu\nu,\lambda} = 0 \quad (26)$$

$\Gamma_\lambda^\sigma$  is a matrix of importance in Schroedinger's theory

It is of interest to examine (26) in the light of (24)

If we write (24) in the form  $\alpha_{\mu,\lambda} = 0$ , we can shorten the process, remembering that

$$\alpha_{\mu,\lambda} = \alpha_{\mu,\lambda} - T_{\mu\lambda}^\sigma \alpha_\sigma = 0 \quad (27)$$

Now

$$2\gamma_{\mu\nu} = \alpha_\mu \alpha_\nu + \alpha_\nu \alpha_\mu,$$

omitting the unit matrix for convenience. Thus

$$2\gamma_{\mu\nu,\lambda} = \alpha_\mu \alpha_{\nu,\lambda} + \alpha_{\nu,\lambda} \alpha_\mu + \alpha_\nu \alpha_{\mu,\lambda} + \alpha_{\mu,\lambda} \alpha_\nu$$

On substituting from (27) the right-hand side becomes

$$T_{\nu\lambda}^\sigma (\alpha_\mu \alpha_\sigma + \alpha_\sigma \alpha_\mu) + T_{\mu\lambda}^\sigma (\alpha_\nu \alpha_\sigma + \alpha_\sigma \alpha_\nu) = 2\gamma_{\mu\sigma} T_{\nu\lambda}^\sigma + 2\gamma_{\nu\sigma} T_{\mu\lambda}^\sigma$$

This vanishes and (26) is preserved if

$$\gamma_{\mu\sigma} T_{\nu\lambda}^\sigma + \gamma_{\nu\sigma} T_{\mu\lambda}^\sigma = 0 \quad (28)$$

With the form (11) for  $T_{\mu\nu}^\sigma$ , this relation holds or the important equation (26) is preserved

The remarks made on the relation between the notation of the earlier paper and Schroedinger's apply to the notation employed here. Certain differences

\* 'Proc Roy Soc., A, vol 141, p 363 (1933)

† 'Sitzber. Preuss. Akad. Wiss., No 24, p. 346 (1932)

between this method and that of Schroedinger's are brought to light in the discussion. We can now see the connection between our theory and that of matrices. Bargmann's suggestion (equation (25)) leads from one to the other.

The relation may be satisfied by writing

$$\Gamma_\lambda = \frac{1}{2} \alpha^\rho \alpha^\sigma T_{\sigma\lambda\rho}, \quad (29)$$

where

$$T_{\sigma\lambda\rho} = \gamma_{\rho\beta} T_{\sigma\lambda}{}^\beta \quad (30)$$

The result follows at once from the matrix relation

$$\alpha_\mu \alpha^\nu + \alpha^\nu \alpha_\mu = 2\delta_\mu^\nu I \quad (31)$$

and from (28), which is equivalent to

$$T_{\nu\lambda\mu} + T_{\mu\lambda\nu} = 0 \quad (32)$$

The further points of contact of the matrix method and that of this paper will be appreciated by a study of Schroedinger's work, any further development here could only be a repetition of it.

The value of  $\Gamma_\lambda$  should be obtained in terms of the  $\Lambda_{\sigma\lambda}{}^\beta$ , since these quantities are made the most fundamental of the theory. The necessary calculation follows at once from (11) and (29). If we make use of the matrix  $\sigma^{\mu\nu}$  given by

$$\alpha^\mu \alpha^\nu + \alpha^\nu \alpha^\mu = 2s^{\mu\nu} \quad (33)$$

we find

$$\Gamma_\lambda = \frac{1}{2} \Lambda_{\lambda\sigma}{}^\beta \delta_\beta{}^\sigma - \frac{1}{4} \gamma_{\lambda\beta} \Lambda_{\sigma\rho}{}^\beta s^{\sigma\rho}, \quad (34)$$

$\delta_\beta{}^\sigma$  is written for  $\delta_\beta{}^\sigma I$  (31)

Schroedinger regards "spur  $\Gamma_\lambda$ " as  $4\phi_\lambda$ , where  $\phi_\lambda$  is the electromagnetic potential.

In our work we have taken

$$\begin{aligned} \text{"Spur } \Gamma_\lambda \text{"} &\propto \Lambda_{\lambda\sigma}{}^\sigma \\ &\propto \frac{1}{\psi} \frac{\partial \psi}{\partial x^\lambda} \end{aligned} \quad (35)$$

This quantity thus occupies a position similar to the electromagnetic potential in the four-dimensional theory. We have previously suggested that the corresponding quantity in the five-dimensional theory is Wilson's vector

$$\pi_\lambda = p_\lambda + \frac{e}{c} \phi_\lambda$$

If this can be extended to the present case we can write

$$\frac{1}{\psi} \frac{\partial \psi}{\partial x^A} \propto \pi_A \quad (36)$$

### *The Invariance of the Quantum Equations*

From the result of the formulation and study of the problem given here, it would seem that the second order equation of the quantum theory in the form (13) is invariant in the system proposed here and the suggestion is that this is the type of invariance which is applicable to the quantum equations.

With regard to the first order equations, it is as if we would pass backward from the wave equation for electromagnetic waves to Maxwell's equations. Their invariant form can be determined by the introduction of matrices and adapting them to fit the notation introduced here.

If we are correct in our suggestion that the second order equation is nothing but the curvature equation then the test will be that having satisfied the notation the first order equations lead to (13).

It would appear that Schroedinger's system satisfies both these requirements, but we suggest that it should be five-dimensional.

### *Summary*

The paper gives a relativistic treatment of the second order equation of the quantum theory. This equation turns out to be the curvature equation in the system proposed, corresponding to the equation of the theory of relativity,  $G_{\mu\nu} = 4\lambda$ . The basis of the work is a modification of the Riemannian' covariant operator and it is shown that the geometric background of the theory permits of some latitude. For this reason an appeal is made to geometrical methods to obtain an indication of the form of the equations but no special theory of parallelism is introduced directly.

The relation to the matrix theory, especially in the form recently developed by Schroedinger is discussed and the two theories are found to be in harmony.

The equation proposed is invariant and throws light on the kind of invariance appropriate to the quantum theory. It is, moreover, an advance on previous equations in that it contains terms appropriate to the gravitational and electromagnetic fields, which come in only with difficulty in other methods.

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possible to decide confidently between the two possibilities on experimental grounds

The arguments used in this paper are based entirely on the need of securing a steady state of electrolysis when a steady current flows through the solution. There must be no time variable concentrations in any part of the solution or on the electrodes. The effective separations found are achieved in such steady states. It is thought that perhaps the full implications of the steady state requirements have not always been realized by workers in this field, and therefore that the attempt, here presented, at a full and careful exposition of them may not be without some general interest.

In NaOH (or other alkaline) solutions the current is usually regarded as being carried by the movement of the ions  $Na^+$  and  $(OH)^-$  both more or less hydrated. We can think of  $(OH)^-$  as moving to the anode and  $Na^+$ , or owing to the hydration  $(NaOH)_2^+$  if we prefer it, as moving to the cathode where it surrenders  $H^+$  and becomes again  $(Na^+ OH^-)$ . This process is formally equivalent to the surrender of an  $H^+$  by an  $OH_2$  water molecule, an  $(OH)^-$  being left behind in the solution free to travel to the anode. That this process of surrender may require the presence of  $Na^+$  before it can occur does not affect its essential form. At the anode  $(OH)^-$  ions become discharged on arrival and oxygen gas  $O_2$  is finally evolved by some such chain of surface reactions as  $2(OH)^- \rightarrow (H_2 + O)$ ,  $2O \rightarrow O_2$  (gas). But as these processes do not evolve hydrogen they are of little interest to us here and we shall not have much occasion to refer to them again. In the next section we shall have occasion to examine more closely the steady state requirements as to the movement of ions, when it will appear, of course, that in the steady electrolysis of water the  $Na^+$  ions on the average do not move and the whole current is carried by the  $(OH)^-$  ions.

We shall in general in this paper discuss directly only alkaline solutions, since it is for such solutions that most is known. But all our arguments apply with obvious slight changes to electrolysis in acids in which the positive  $(OH_2)^+$  ions alone move and convey the current.

§ 2 *General Description of the various Processes assumed to occur*—The arguments which we shall apply will often be general enough not to need any detailed specification of the mechanism by which a particular process is occurring. It is none the less necessary for clarity to specify the broad outlines of the processes occurring at and near the cathode, to which almost any special mechanism must conform. It is therefore convenient to start by distinguishing three phases: (1) the interior of the solution generally, (2) its boundary layer

in contact with the cathode, and (3) the surface of the cathode itself. When, if ever, a valid distinction is to be drawn between (1) and (2) is a matter for investigation (§ 5). When a steady current flows we must consider the exchanges of the two hydrogens between these three phases and the evolved gas. No accumulations can form anywhere except in the evolved gas which is ultimately formed at the expense only of the general body of hydrogen in the water of the interior of the solution.

From the boundary layer of the solution hydrogen is fed on to the surface of the cathode. This is an essential part of the current-carrying process. It will be necessary to consider whether or not the only exchanges of hydrogen between the cathode and the boundary layer are the current-carrying depositions of hydrogen atoms, or whether important balanced exchanges take place in addition. Finally since the hydrogen is certainly supplied to the cathode surface phase as atoms by the above process and evolved into the gas phase as molecules, it is necessary to consider a final process in the evolution of hydrogen gas in which atoms of hydrogen combine to form molecules on the cathode surface.

§ 3 *Passage of Hydrogen from the Boundary Layer to the Cathode*—As we have said in the introductory section the essential action on the cathode which enables the current to flow is the break up of a water molecule  $\text{OH}_2$  which surrenders a proton  $\text{H}^+$  to the cathode and leaves an extra  $(\text{OH})^-$  ion behind in the water free to move to the anode. It is a matter of comparative indifference whether or not the action involves intimately the presence of  $\text{Na}^+$  so as to be more accurately described as



This process has been visualized by Gurney rather as a process of discharge of the ion  $(\text{NaOH}_2)^+$  by an electron jump from the cathode followed by a process of ejection of the unstable extra H atom from  $\text{NaOH}_2$  which then becomes  $\text{Na}^+ (\text{OH})^-$  while the H atom combines with the metal surface. (The word "combine" is here used loosely.) Whatever the correct version may be, it is agreed universally that the process is atomic and that H not  $\text{H}_2$  is supplied to the cathode.

If  $n_1^b$ ,  $n_2^b$  are the concentrations of  $\text{H}^1$  and  $\text{H}^2$  in the cathode boundary layer, then as the process is atomic and a first order reaction we may assume that  $\text{H}^1$  atoms are deposited at a rate  $\kappa_1 n_1^b$  and  $\text{H}^2$  atoms at a rate  $\kappa_2 n_2^b$ , where  $\kappa_1$ ,  $\kappa_2$  are independent of  $n_1^b$ ,  $n_2^b$ . It is not necessarily true that  $\kappa_1$  equals  $\kappa_2$ . The  $\kappa$ 's might depend on the amount of hydrogen already present on the

surface, on the temperature, and on the voltage drop from the cathode boundary layer to the cathode (the over-potential) So far as we know at present the concentrations of the hydrogens which are in a state to be transferred to the electrode, because, for example, they are attached to positive ions will be the same as their concentrations on the complete boundary layer

If the  $\kappa$ 's depend on the hydrogen already present on the cathode, let  $n_1'$  and  $n_2'$  be the concentrations of the two sorts of hydrogen in the cathode surface Then the only dependence of the  $\kappa$ 's on  $n_1'$  and  $n_2'$  which is at all likely is that the  $\kappa$ 's are of the form

$$\kappa_1\{1 - \gamma(n_1' + n_2')\}, \quad \kappa_2\{1 - \gamma(n_1' + n_2')\}$$

so that their ratio would be unaffected This dependence would occur if the hydrogen to be deposited required a vacant metal atom to combine with It would be indifferent to prior occupation by  $H^1$  or by  $H^2$

If the current carrying process is the only means by which hydrogen is conveyed from the solution or rather its boundary layer to the cathode then the whole exchange will be at the rates  $\kappa_1 n_1^b$ ,  $\kappa_2 n_2^b$  and all in the one direction,  $\kappa_1 n_1^b + \kappa_2 n_2^b$  will be fixed by the current density It is, however, uncertain whether this does constitute the major part of the hydrogen exchange There is some evidence that even when no current is flowing there may be some considerable balanced exchange of hydrogen between the electrode and the water, particularly the ions of the boundary layer We shall then have an equilibrium state controlled by the equations,

$$\kappa_1 n_1^b = \mu_1 n_1', \quad \kappa_2 n_2^b = \mu_2 n_2',$$

where  $\mu_1 n_1'$ ,  $\mu_2 n_2'$  are the rates of transfer from the electrode back to the solution, and  $\mu_1$ ,  $\mu_2$  are independent of the  $n$ 's In this case all the rates may be large compared with the rates demanded by any ordinary current density and then these equilibrium conditions will not be seriously disturbed by the passage of the current

The two conditions detailed above are, of course, the two possible limiting cases and every intermediate case is possible We shall, however, for simplicity only discuss these two limits which are sufficient for illustration

§ 4 *The Evolution of Hydrogen Gas at the Cathode*—The process of formation of  $H_2$  gas appears to be one which takes place actually on the surface or in the immediate surface layer of the metal—in a metallic phase, that is, which is rich in adsorbed or absorbed H atoms This process must clearly be assumed to be a second order reaction The relevant concentrations of the  $H^1$  and  $H^2$

atoms will be  $n_1^e$  and  $n_2^e$  as already defined. [Conceivably the relevant concentrations might differ from these values, but if so they could only differ by factors which can be ignored because they could be absorbed in the  $\lambda$ 's shortly to be defined.] Molecules  $\text{H}^+\text{H}^+$ ,  $\text{H}^+\text{H}^e$  and  $\text{H}^e\text{H}^e$  may therefore be assumed to be formed respectively at rates

$$\lambda_{11} (n_1^e)^2, \quad \lambda_{12} n_1^e n_2^e, \quad \lambda_{22} (n_2^e)^2$$

where the  $\lambda$ 's are independent of the  $n$ 's.

At this stage again there appear *a priori* to be two possible limiting cases though now they do not lead to distinct forms. There may be an equilibrium concentration of hydrogen molecules in and on the metal surface which is practically zero so that effectively all the molecules formed immediately escape as gas. In this case the rate of removal of  $\text{H}^+$  atoms from the water will be

$$2\lambda_{11} (n_1^e)^2 + \lambda_{12} n_1^e n_2^e, \quad (2)$$

and the rate of removal of  $\text{H}^e$  atoms

$$2\lambda_{22} (n_2^e)^2 + \lambda_{12} n_1^e n_2^e. \quad (3)$$

The  $\lambda$ 's need not be equal.

On the other hand, there may be a significant equilibrium concentration of molecules in the metal maintained by rates of formation and destruction sufficiently rapid for the evolution of gas required by the current transport not to disturb it. In that case we must obviously expect that the rates of evolution of  $\text{H}_2$  molecules of each type will be proportional to their equilibrium concentrations in the metal surface  $n_{11}^e$ ,  $n_{12}^e$  and  $n_{22}^e$  respectively.

The rate of removal of  $\text{H}^+$  atoms from the water will then be

$$2v_{11} n_{11}^e + v_{12} n_{12}^e,$$

and the rate of removal of  $\text{H}^e$  atoms

$$2v_{22} n_{22}^e + v_{12} n_{12}^e.$$

Since, however, we must expect equilibrium relationships between the various  $n$ 's similar to those of a perfect gas, and of the forms

$$n_{11}^e = f_{11} (n_1^e)^2, \quad n_{12}^e = f_{12} n_1^e n_2^e, \quad n_{22}^e = f_{22} (n_2^e)^2,$$

where the  $f$ 's are independent of the  $n$ 's, the formulæ of this limiting case reduce to (2) and (3). Presumably, therefore, it is not necessary to distinguish here between these limits, or any of the possible intermediate cases.

§ 5 *Transport of Hydrogen to and fro between the Cathode Boundary Layer and the Interior of the Solution*—In any steady state the  $Na^+$  ions may not accumulate and therefore cannot be moving, so that the whole current is carried by the movement of the  $(OH)^-$  ions to the anode. This movement "conveys" hydrogen to the cathode by allowing an  $(OH)^-$  ion to be replaced by an  $OH_2$  molecule. Besides this transport the process of diffusion (molecular and turbulent) conveys hydrogen (in the water molecules and ions) to and fro between the boundary and the interior.

If all diffusion could be neglected, since there are no accumulations anywhere in the boundary region the two hydrogens would be evolved (no matter what the mechanism) at rates in the ratio of the currents carried by the two types of hydrogens—that is, in the ratio

$$E_1 : E_2 = pD_1 : D_2, \quad (4)$$

where  $p$  is the ratio of the mobilities of the two hydrogens. The yield of each must equal the supply.<sup>1</sup> The mobility of  $H^1$  means here the rate of effective movement due to the replacement of the grouping  $(OHH^1)$   $(OH)^-$  by  $(OH)(OHH^1)$ , where the unindexed  $H$  may be either  $H^1$  or  $H^2$ , with a similar meaning for the mobility of  $H^2$ . The factor  $p$  is probably\* about 1.5, but we shall not stay to discuss it more closely, since it can be shown that diffusion will always mask the current transport and keep the hydrogen concentrations in the boundary layer effectively in the ratio  $D_1 : D_2$ , which is that of the water as a whole. We now proceed to consider this effect of diffusion.

Let  $n_1$  and  $n_2$  be the total concentration of the two hydrogens at any point in the water—hydrogen nuclei to be precise. Then since the number of water ions can be locally variable it is not necessary that  $n_1 + n_2 = \text{const}$ . They are independently variable. Under the influence of a given field let the number of  $H^1$  and  $H^2$  nuclei which cross unit area in unit time be  $\alpha_1 n_1$  and  $\alpha_2 n_2$  respectively. The  $\alpha$ 's are proportional to the mean velocities of drift of the two nuclei and independent of the  $n$ 's, but they are not *mobilities* since  $n_1$  and  $n_2$  are total molecular not ionic concentrations. When any concentration gradient is set up, along the  $x$  axis for example, the numbers of  $H^1$  and  $H^2$  nuclei which diffuse back per  $cm^2$  per second owing to the self diffusion of the water molecules, are respectively

$$-d_1 \frac{\partial n_1}{\partial x}, \quad -d_2 \frac{\partial n_2}{\partial x}$$

\* See Lewis and Doody, 'J. Amer. Chem. Soc.', vol. 55, p. 3504 (1933) for measurements on the replacement  $(OH_2)^+ OH_2$  by  $(OH_2)(OH_2)^+$ . For a theory which allows for these mobilities to be different for  $H^1$  and  $H^2$  see Bernal and Fowler ('J. Chem. Phys.', vol. 1, p. 515 (1933)).

The  $d$ 's are probably nearly equal, and in order of magnitude equal to the coefficient of self-diffusion of water. We may suppose that the whole flow takes place along the  $x$  axis. Then since no accumulations are occurring

$$\alpha_1 n_1 - d_1 \frac{\partial n_1}{\partial x} = E_1, \quad \alpha_2 n_2 - d_2 \frac{\partial n_2}{\partial x} = E_2 \quad (5)$$

The equations hold for values of  $x$  from the cathode ( $x = 0$ ) so far into the solution that the water between such value of  $x$  and the cathode is an appreciable fraction of the whole solution. Since the field gradients are not constant owing to space charge the  $\alpha$ 's strictly depend on  $x$ , but this can be ignored here. We return to it in § 8.

Equations (5) can be solved at once and give

$$n_1 = \frac{E_1}{\alpha_1} + C_1 e^{\alpha_1 x/d_1}, \quad n_2 = \frac{E_2}{\alpha_2} + C_2 e^{\alpha_2 x/d_2} \quad (6)$$

The constants  $C_1$  and  $C_2$  must be adjusted to give the correct boundary values to  $n_1$  and  $n_2$  at  $x = 0$  required to transfer the observed quantities  $E_1$  and  $E_2$  of evolved gas. These boundary values do depend, therefore, on the cathode mechanism. It is clear at once that if  $\alpha_1/d_1$  is so large that while formulæ (6) still apply  $\alpha_1 x/d_1$  becomes effectively  $-\infty$ , then since at the same time  $n_1/n_2 \rightarrow D_1/D_2$ , we have  $E_1/E_2 = (\alpha_1/\alpha_2) D_1/D_2 = p D_1/D_2$  as already stated. If, however,  $\alpha_1/d_1$  is so small that  $\alpha_1 x/d_1 \approx 0$  throughout the region of validity of (6) then there is perfect mixing by diffusion in spite of the current and  $n_1/n_2 = n_1^0/n_2^0$  everywhere, so that

$$n_1^0/n_2^0 = D_1/D_2 \quad (7)$$

Everything then depends on the numerical order of  $\alpha l/d$  where  $l$  is the scale of the cell, say, the distance between the electrodes. The numerical value of  $\alpha$  can, of course, be derived from the mobility, but it is more simply derived by observing that  $\alpha_1 n_1$  is approximately equal to  $E_1$ . A current of 1 ampere/cm<sup>2</sup> requires a net flow of  $6 \times 10^{18}$  protons per cm<sup>2</sup> per second. Since  $n_1$  is of the order  $7 \times 10^{22}$  protons per cm<sup>3</sup>  $\alpha_1 \approx 10^{-4}$ , and  $\alpha_2$  is of the same order.

The numerical value of  $d_1$  or  $d_2$  is not yet known by direct observation of the molecular diffusion of heavy water into ordinary water, but its order can probably be safely derived from the known values for the diffusion coefficient for the diffusion of other small molecules into water. The values of all such coefficients of diffusion are of the order  $10^{-3}$  cm.<sup>2</sup>/sec. Thus

$$\alpha l/d \approx 10^{-7}.$$

For smaller current densities the coefficient is proportionately smaller. Thus, except for obviously large scale apparatus molecular diffusion is capable of keeping the boundary concentrations normal for current densities up to 10 amp/cm.<sup>2</sup>, which are greater than any hitherto used.

Theoretically relation (7) might fail for very great current densities, but these are so great that they would induce other and more efficient means of mixing by turbulence due to the formation of bubbles on the cathode. It is therefore doubtful if (7) ever fails.

§ 6 *The Equations of Steady (Alkaline) Electrolysis*—Having thus cleared the ground we can write down the general equations of electrolysis to which the process will conform no matter what its detailed mechanism. When the process is steady any differential coefficients such as  $dn/dt$  must vanish. The concentrations of the two hydrogens in the boundary layer are  $D_1$ ,  $D_2$  for the reasons given in § 5. These equations are

$$\frac{dn_1}{dt} = \kappa_1 D_1 - \mu_1 n_1 - E_1 = 0 \quad (8)$$

$$\frac{dn_2}{dt} = \kappa_2 D_2 - \mu_2 n_2 - E_2 = 0, \quad (9)$$

$$E_1 = 2\lambda_{11}(n_1)^2 + \lambda_{12}n_1 n_2, \quad (10)$$

$$E_2 = 2\lambda_{22}(n_2)^2 + \lambda_{12}n_1 n_2. \quad (11)$$

There are some important special cases

*Case (1)*—*The to and fro exchange of hydrogen between the solution and the electrode small compared with  $E_1$ ,  $E_2$* . Under these conditions  $\mu_1 n_1$  and  $\mu_2 n_2$  can be neglected in (8) and (9), and we find simply no matter what values the  $\lambda$ 's have

$$\frac{E_1}{E_2} = \frac{\kappa_1 D_1}{\kappa_2 D_2} \quad (12)$$

This specifies an evolution of gas of the observed form if under the best working conditions  $\kappa_1/\kappa_2 \approx 6$ . This case provides, of course, the explanation proposed by Polanyi for the observed separations. The  $\kappa$ 's could have such a ratio if the controlling mechanism of the deposition of hydrogen on the cathode is the passage of the hydrogen nuclei through a small potential barrier by the tunnel effect when the rate of penetration of  $H^1$  for a given concentration can easily exceed that of  $H^2$  by a factor of the order of 10. Gurney's theory of the hydrogen over-potential would then be inapplicable to the phenomenon.\*

[\* *Note added February 20th*—The  $\kappa$ 's could still have such a ratio on Gurney's theory. See Bell and Wolfenden, 'Nature,' vol 133, p 25 (1934).]

It is, however, not certain that the conditions of case (i) are fulfilled. It is known from the observations of Polanyi\* himself, stimulated by observations of Oliphant,† that a very considerable exchange of hydrogen can take place between water and hydrogen gas in contact with a suitable catalysing surface in Polanyi's case platinum black. This exchange, as is pointed out, appears to be precisely that with which we are concerned here, but its rate can hardly be estimated from what is yet known. The action might be of one or both of two kinds: (a) a simultaneous interchange of two hydrogen nuclei, one in a water molecule and one adsorbed on the surface by a process of the type  $\text{OHH}' + \text{H}''\text{Me} \rightarrow \text{OHH}'' + \text{H}'\text{Me}$ , (b) a continual formation and discharge of water ions in the neighbourhood of the catalytic surface leading to balanced reactions of the types (i)  $(\text{OH}_3)^+ + \text{Me}^- \rightleftharpoons \text{OH}_2 + \text{HMe}$  and (ii)  $(\text{OH})^- + \text{HMe}^+ \rightleftharpoons \text{OH}_2 + \text{Me}$ . It is, therefore, important to discuss equations (8)–(11) in other conditions than those of case (i).

Case (ii) — *The to and fro exchange of hydrogen between the solution and the electrode large compared with  $E_1$  and  $E_2$* . Under these conditions  $E_1$  and  $E_2$  can be neglected in (8) and (9). We, therefore, obtain from them

$$n_1 e = f_1 D_1, \quad n_2 e = f_2 D_2$$

The coefficients  $f_1$  and  $f_2$  may well be functions of the over-potential, that is, the voltage drop at the cathode surface required to carry a given current density. It may, therefore, be well to write them sometimes as  $f_1(V)$ ,  $f_2(V)$  or as  $f_1(I)$ ,  $f_2(I)$  to recall this probable dependence. Equations (10) and (11) then give

$$\frac{E_1}{E_2} = \frac{2\lambda_{11}f_1^2D_1 + \lambda_{12}f_1f_2D_1D_2}{2\lambda_{22}f_2^2D_2 + \lambda_{12}f_1f_2D_1D_2} \quad (13)$$

This equation cannot be of the observed form unless it reduces to  $E_1/E_2 = \alpha D_1/D_2$ . It can, however, so reduce if

$$\frac{2\lambda_{11}}{\lambda_{12}} - \frac{\lambda_{11}}{2\lambda_{22}} = q, \quad (14)$$

when

$$\frac{E_1}{E_2} = q \frac{f_1 D_1}{f_2 D_2} \quad (15)$$

This equation will, of course, represent the results of the best experiments with nickel electrodes and currents of the order 1 amp/cm.<sup>2</sup> if then  $qf_1/f_2 = 6$

\* Horvut and Polanyi, 'Nature,' vol 132, p 819 (1933)

† 'Nature,' vol 132, p 675 (1933)



Both  $q$  and the  $f$ 's should depend on the temperature, but  $q$  should not depend on the voltage and therefore not on the current density

At first sight it is hardly to be expected that there is any reason why (14) should be satisfied—even approximately. It requires the relationship  $\lambda_{11}\lambda_{22}/\lambda_{12}^2 = \frac{1}{4}$  to hold between the  $\lambda$ 's, but there is after all some *a priori* reason to expect such a connection.

Let us, for example, take that view of the mechanism of gas evolution which regards the  $\lambda$ 's as collision recombination rates for the formation of  $H_2$  molecules on the metal surface. Let us then consider the  $H$  atoms on the surface as an ideal quasi-gaseous mixture of atoms  $H^1$  and  $H^2$  of concentrations  $n_1^*$  and  $n_2^*$ . The total numbers of effective collisions have been defined as  $\lambda_{11}(n_1^*)^2$ ,  $\lambda_{12}n_1^*n_2^*$  and  $\lambda_{22}(n_2^*)^2$ . If these were the total number of collisions of freely moving gas atoms then

$$\lambda_{11} \propto \frac{1}{2}\bar{C}_{11}, \quad \lambda_{12} \propto \bar{C}_{12}, \quad \lambda_{22} \propto \frac{1}{2}\bar{C}_{22},$$

where the  $C$ 's are the mean relative velocities of the two sorts of atom. If there is an excitation energy required for the success of the recombination then the number of favourable collisions might be proportional respectively to

$$\frac{1}{2}\bar{C}_{11}e^{-x_{11}/kT}, \quad \bar{C}_{12}e^{-x_{12}/kT}, \quad \frac{1}{2}\bar{C}_{22}e^{-x_{22}/kT}.$$

But

$$\bar{C}_{11} \bar{C}_{12} \bar{C}_{22} = \sqrt{\left\{\frac{1}{m_1} + \frac{1}{m_1}\right\}} \sqrt{\left\{\frac{1}{m_1} + \frac{1}{m_2}\right\}} \sqrt{\left\{\frac{1}{m_2} + \frac{1}{m_2}\right\}}$$

Hence

$$\frac{\lambda_{11}\lambda_{22}}{\lambda_{12}^2} = \frac{1}{4} \frac{2(m_1m_2)^{\frac{1}{2}}}{m_1 + m_2} e^{-(x_{11}+x_{22}-x_{12})/kT} \quad (16)$$

Now it can well happen that  $x_{11} + x_{22} - 2x_{12} \geq 0$ . For though the activation energy required by  $H^2H^2$  may well be substantially greater than that required by  $H^1H^1$ , in that case the energy required for  $H^1H^2$  is likely to be very nearly the mean value of the other two. The mass factor is  $2\sqrt{2/3} = 0.94$  even for masses so different as those of  $H^1$  and  $H^2$ . Thus  $\lambda_{11}\lambda_{22}/\lambda_{12}^2$  does not on this view seem likely to deviate much from  $\frac{1}{4}$  and may well be almost exactly equal to it, even if  $2\lambda_{11}/\lambda_{12} = q$  is distinctly different from unity. A similar close equality between  $\lambda_{11}\lambda_{22}/\lambda_{12}^2$  and  $\frac{1}{4}$  can be shown to be expected if we take the other view of the mechanism in which the  $\lambda$ 's represent velocities of escape of already formed molecules, and account is taken of the equilibrium constants of the dissociating quasi-gas on the metal surface.

In this case we see that it is easily possible that the whole differential factor in the rates of evolution of  $H^1$  and  $H^2$  may come from preferential molecule

formation on the electrode surface—the factor  $f_1/f_2$  being then close to unity, but with separation still possible owing to the factor  $q$

It is not maintained here that case (ii) represents a true view of the electrode mechanism—only that it is at present a possible one which is not ruled out by any experiments yet reported. It is important to note that if it were a true view then Gurney's explanation of the hydrogen over-potential might still stand.

There is, however, one piece of evidence in this field which may possibly if correctly interpreted, lead to a discrimination against Gurney's version, or rather against any theory which allows  $\kappa_1 \simeq \kappa_2$  and casts the onus for separation on the surface recombination—that is, the apparently reduced efficiency of the separation at lower current densities or at lower over-potentials. It seems possible that this means that at the higher current densities we have case (i) fully developed, but at lower current densities the to and fro exchanges are no longer negligible by comparison so that we tend to move towards the conditions of case (ii). It does not seem possible to refer this observed variation directly to variations of  $\kappa_1/\kappa_2$  with over potential, for in that case the higher the over-potential the more nearer equal one must expect the  $\kappa$ 's to become and the efficiency should be least at high current densities where it appears to be greatest. Since this explanation may be important we consider a third special case which might apply to it.

*Case (iii).—Neither the  $\mu$ 's nor the  $k$ 's negligible in (8) and (9) but  $\lambda_{11}\lambda_{22}/\lambda_{12}^2 = \frac{1}{4}$ .* These are the simplest type of conditions intermediate between those of cases (i) and (ii). We find

$$\frac{E_1}{E_2} = \frac{2\lambda_{11}\mu_2}{\lambda_{12}\mu_1} \frac{\kappa_1 D_1 - E_1}{\kappa_2 D_2 - E_2} \quad (17)$$

It may with the help of this equation be possible to trace a transition between the two limits in which

$$\frac{E_1}{E_2} = \frac{2\lambda_{11}f_1}{\lambda_{12}f_2} \frac{D_1}{D_2} \quad \frac{\kappa_1 D_1}{\kappa_2 D_2}$$

for low and high current densities respectively. It must be remembered that the  $\kappa$ 's at least are functions of the over-voltage and therefore of the current or  $E_1 + E_2$ .

§ 7 *Conclusions*—We may summarize this discussion as follows—

(1) If the transport of hydrogen by the current could overwhelm the transport by the natural processes of diffusion the rates of evolution of  $H^+$  and  $H^2$  would be in the ratio of the mobilities of the two hydrogen ions no matter

what the electrode mechanism. This condition, however, is never fulfilled in practice and diffusion is always ample to keep the two hydrogens in normal concentration ratios on the cathode boundary

(2) If the reversible exchange of hydrogen atoms between the cathode surface and the boundary layer of the liquid can be neglected compared with the hydrogen deposited by the current,  $i$ , at sufficiently high current densities the ratio of the rates of evolution will be of the form  $\kappa_1 D_1 / \kappa_2 D_2$ . This form is in agreement with observation

(3) If the reversible exchange of hydrogen atoms between the cathode surface and the boundary layer of the liquid is fast compared with the hydrogen deposited by the current— $i$ , at sufficiently low current densities—then equilibrium concentrations of the hydrogens on the cathode will be in the ratio  $f_1 D_1 / f_2 D_2$ , where  $f_1 / f_2$  will in general depend on the over potential. The ratio of the rates of evolution of the hydrogen atoms will be  $q f_1 D_1 / f_2 D_2$  if a certain natural condition is satisfied so that the separation may be effected by  $q$  ( $f_1 / f_2 \geq 1$ ) which is a factor arising from the relative efficiency of  $H^{11}H^1$  and  $H^{11}H^2$  collisions in forming molecules, or it may be effected by  $f_1 / f_2$  ( $q \leq 1$ ) and so be a result of different over-voltages for the two hydrogens

(4) It is not possible yet to conclude that the over-potential factors  $\kappa_1 / \kappa_2$  or  $f_1 / f_2$  are necessarily the separating agency. Thus we cannot immediately rule out Gurney's theory of the hydrogen electrode. There is, however, this factor in favour of throwing the onus for the best separation factor (about 6) on to  $\kappa_1 / \kappa_2$ —that the separation appears to fall off at lower current densities. If we refer the high current high efficiency to  $\kappa_1 / \kappa_2$ , we can refer the lowering at lower currents to a change over towards  $q f_1 / f_2$ .

§ 8 *The Equations of Steady Electrolysis in an Alkaline Solution*—In this section we are no longer concerned with the separation of the two hydrogens, but strictly speaking consider only the way in which the current is conveyed through an alkaline electrolyte when the electrolysis of pure water (either light or heavy) is proceeding steadily. It appears that, familiar as this problem is, something more can be said about the theory of it, by bringing in the ideas of space charge used so successfully in the theory of electron currents in high vacua.

Suppose that the current is flowing everywhere parallel to the  $x$ -axis between plane parallel electrodes. In the solution there are  $Na^+$  ions (or other positives) which on the whole cannot move and therefore have the sole function of creating a space charge  $en_+$ , where  $n_+$  is their concentration per unit volume. Let  $V$  be the electrostatic potential in the solution and let the cathode surface

be  $x=0$  and  $V(0)=0$ . Then since the state of the positive ions is an equilibrium state in a field of potential  $V(x)$ ,

$$n_+ = n_+^0 e^{-eV/RT} \quad (18)$$

The  $(OH)^-$  ions have a concentration  $n^-$ , produce a space charge  $-en^-$  and carry the current by their movement. If  $\alpha$  is their effective mobility under unit field strength then  $\alpha e \partial V / \partial x$  is their average velocity and the current density at any point in the fluid is  $-\alpha e^2 n^- \partial V / \partial x$  in the direction of  $V$  increasing. We may call this current  $I$  (since the conventional current flows in the direction of  $V$  decreasing) so that

$$I = \alpha e^2 n^- \frac{\partial V}{\partial x} \quad (19)$$

The potential  $V$  satisfies Poisson's equation,

$$\frac{d^2 V}{dx^2} = -\frac{4\pi e}{K} (n^+ - n^-), \quad (20)$$

where  $K$  is the dielectric constant of water assumed independent of  $x$ . This is the only place in these equations where  $K$  occurs. Then combining (18), (19) and (20) we obtain an equation which can be integrated once giving

$$\left(\frac{dV}{dx}\right)^2 - \left(\frac{dV}{dx}\right)_0^2 = -\frac{8\pi kT}{K} n_+^0 \{1 - e^{-eV/RT}\} + \frac{8\pi}{K} \frac{1}{\alpha} x \quad (21)$$

We have here assumed that  $V$  and  $\partial V / \partial x$  are continuous functions of  $x$  and that in particular  $V(0)=0$ , the potential of the cathode. The value of  $(\partial V / \partial x)_0$ , on this version of the theory, must represent the over-potential, which here appears not as a discontinuity but as a steep initial potential gradient. On any theory of the cathode mechanism there will be a relationship of the form

$$\left(\frac{dV}{dx}\right)_0 = f(I), \quad (22)$$

or possibly more significantly of the form

$$\left(\frac{dV}{dx}\right)_0 = f(n_+^0, I) \quad (23)$$

This is one of the boundary conditions of the problem.

A second integration in finite terms is not possible. But if we put

$$eV/RT = \phi$$

equation (21) becomes

$$\left(\frac{d\phi}{dx}\right)^2 - \left(\frac{d\phi}{dx}\right)_0^2 = -\frac{8\pi\epsilon^2 n_+^0}{KkT} \{1 - e^{-\phi}\} + \frac{8\pi I\epsilon}{\alpha Kk^2 T^2} x \quad (24)$$

If now we put

$$x = \mu\xi \quad \mu = \left(\frac{KkT}{8\pi\epsilon^2 n_+^0}\right)',$$

then

$$\left(\frac{d\phi}{d\xi}\right)^2 - \left(\frac{d\phi}{d\xi}\right)_0^2 = -(1 - e^{-\phi}) + \lambda\xi, \quad (25)$$

where

$$\lambda = \frac{K^{\frac{1}{2}} I}{(8\pi kT)^{\frac{1}{2}} \alpha \epsilon^2 (n_+^0)^{3/2}} \quad (26)$$

Equation (25) depends on the two parameters  $(\partial\phi/\partial\xi)_0$  and  $\lambda$  and it is a not impossibly laborious task to compute its solutions numerically. We could thus determine

$$\phi = \phi(\xi, \lambda, \left(\frac{\partial\phi}{\partial\xi}\right)_0), \quad (27)$$

and the final boundary condition which has to be applied is to make  $V$  take the correct value on the other electrode. Thus (27) really means

$$V(l) = V_0 = \Phi(l, I, n_+^0, (\partial V/\partial x)_0) \quad (28)$$

Again from equation (27) since  $\phi$  is known  $n_+^0$  can be determined in terms of the total concentration of electrolyte in the whole cell, which is fixed. Thus (23) and (28) together give the current voltage relationship of the cell and (27) the distribution of potential, from which the other distributions follow. We have apparently not satisfied a second boundary condition analogous to (23) at the anode, namely,

$$\left(\frac{dV}{dx}\right)_l = g(n_+^0, I),$$

but the state of the anode is not fixed since the number of available  $(OH)^-$  ions is not fixed, and therefore the anode can and must adjust itself until it can accept a current of the proper density from the boundary layer of the solution in its determined state. It is only by a crude approximation that these molar equations can be used right down to the boundary layer itself. But there is this essential difference between the requirements of the two electrodes when one of the ions is present in fixed amount and the other is indefinite.

Further study of the equations of this section will be undertaken.

*Summary*

This paper discusses in a general way the theory of the separation of hydrogen and deuterium by the electrolysis of water. It is an amplified account of remarks made at the recent discussion on heavy hydrogen and included in the published report. In discussing the theory of this separation, it is necessary to remember that the process is a steady one and that no accumulations of any substance are going on anywhere except in the evolved gas where hydrogen and deuterium accumulate at the expense of the general water supply. The requirements of a steady state are systematically applied to the various possible mechanisms of the hydrogen electrode which have been suggested. It is shown that the self diffusion of water is important and suffices to keep the  $H/D$  ratio near the cathode normal in spite of the different mobilities of the hydrogen and deuterium ions. It is next shown that if there were no to and fro exchange of hydrogen between the solution and the cathode, then the separation would depend on differential rates of transition for protons and deuterons according to the suggestions put forward by Polanyi, but that this exchange certainly exists so that Polanyi's explanation, though still possible, is no longer unique. The conditions for this more general state of affairs are then discussed, and it is shown that so far as existing evidence goes, the separation may be due either to different over-potentials as proposed by Polanyi, or to different combination rates for molecule formation on the surface, or partly to one and partly to the other. The paper concludes with a formulation of the general space charge equations which must govern a steady electrolytic current in the alkaline electrolysis of water.

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## *Experiments on Heavy Hydrogen \*—Part I*

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Cambridge

(Communicated by E. H. Rideal, F.R.S.—Received January 29, 1934)

### *Method of Analysis*

With the exception of those investigations in which spectroscopic† or mass spectrographic‡ methods were applicable, earlier workers on the heavy hydrogen isotope (deuterium or diplogen denoted by  $H^2$  or by D) have determined the ratio H to D from measurements of the specific gravity§ of the water obtained from the heavy hydrogen. One per cent of diplogen causes an increase of about one part in 1000 in the specific gravity. The density measurement may be made with great accuracy, but at least 3.4 mg. of water are required for every determination||

A convenient and reliable determination can be made with a very much smaller quantity of material, both accurately and rapidly, utilizing the thermal conductivity of the gas to indicate its composition in the following manner¶

In fig. 1 the specific heats of normal  $H_2$ ,  $D_2$ , and HD are plotted against the absolute temperature. The rotational specific heats were calculated according to the general formula\*\*

$$C_{rot} = -R \frac{d}{dT} \frac{d \ln Q}{d(1/T)}, \quad Q = \sum p_i e^{-\frac{\epsilon_i}{kT}}$$

\* 'Nature,' vol. 132, p. 894 (1933)

† Urey, Brickwedde, and Murphy, 'Phys. Rev.,' vol. 40, p. 1 (1932)

‡ Bleakney, 'Phys. Rev.,' vol. 41, p. 32 (1932), Bleakney and Gould, *ibid.*, vol. 44, p. 365 (1933)

§ Lewis and Macdonald, 'J. Chem. Phys.,' vol. 1, p. 341 (1933), Washburn, Smith, and Frandsen, 'Bur. Stand. J. Res.,' vol. 11, p. 453 (1933). More recently a refractive index method has been worked out by Lewis and D. B. Luten, junr ('J. Amer. Chem. Soc.,' vol. 55, p. 5061 (1933)) and by Crist, Murphy and Urey ('J. Amer. Chem. Soc.,' vol. 55, p. 5060 (1933))

|| Giffillan and Polanyi, 'Z. phys. Chem.,' A, vol. 166, p. 255 (1933)

¶ Cf. A. Farkas, 'Z. phys. Chem.,' B, vol. 22, p. 344 (1933). Evidently the normal thermal conductivity method as developed by Schleiermacher and applied by Bonhoeffer and Harteck to their experiments on ortho and para hydrogen ('Z. phys. Chem.,' B, vol. 4, p. 113 (1929)) may be used, but over a thousand times the quantity of gas would be required for each measurement, than in the micro-method.

\*\* See, for example, Glaue, 'J. Amer. Chem. Soc.,' vol. 52, pp. 4908, 4916 (1930)

where  $p_j$  is the quantum weight of the rotational state with the quantum number  $j$  and  $J$  is the moment of inertia. For  $H_2$  and  $D_2$  the corresponding expressions of Dennison\* were used which treat hydrogen as a mixture of the ortho- and para-modifications ( $H_2 = \frac{3}{4}$  ortho  $H_2 + \frac{1}{4}$  para  $H_2$ ,  $D_2 = \frac{2}{3}$  ortho  $D_2 + \frac{1}{3}$  para- $D_2$ ) (see Part II). For the moments of inertia  $J$  of  $D_2$  and  $HD$  the values given by Urey and Rittenberg† were employed.

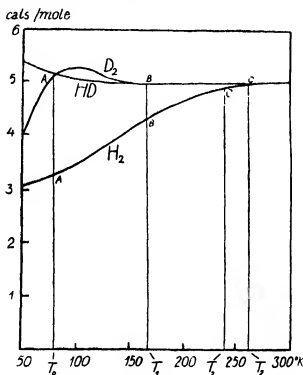


FIG. 1

The heat lost by an electrically heated wire stretched in a narrow tube containing the gas under a low pressure owing to the thermal conductivity of the gas is proportional to the expression  $\int_{T_0}^{T_1} C_d T$ , or on fig. 1 to the area  $T_0 T_1 BA$ , where  $T_0$  is the temperature of the surrounding walls of the vessel and  $T_1$  the temperature to which the wire has been heated by a definite current  $i_1$ . If now the current be increased to  $i_2$  so that the heat given up by the wire to the gas is, say, twice as great as previously the temperature of the wire rises to  $T_2$ .

\* 'Proc. Roy. Soc. A,' vol. 115, p. 483 (1927).

† 'J. Chem. Phys.,' vol. 1, p. 137 (1933).



If the hydrogen in the tube be now replaced by diplogen and the pressure of the gas so regulated that the wire returns to the original temperature  $T_1$  for the current  $i_1$ , this pressure is clearly given by

$$P_{D_1} = P_{H_1} \frac{T_0 T_1 AB}{T_0 T_1 A'B'}$$

but when the current  $i_2$  is again switched on to the wire the temperature attained is now no longer  $T_2$  but  $T'_2$ , where

$$\frac{T_0 T_1 BA}{T_0 T_2 CA} \approx \frac{T_0 T_1 B'A'}{T_0 T_2 C'A'} *$$

From the temperature difference  $T'_2 - T_2$  the D content of the gas may be determined from a calibration curve. In our experimental arrangement (0.01 mm diameter platinum wire, 5 cm long in gas at a pressure of ca 0.04 mm)  $T'_2 - T_2$  for the pair of gases hydrogen and diplogen is about  $20^\circ$  and corresponds to a difference in resistance of about 6 ohms. From fig 1 it is seen that the temperature variation of the specific heat of HD is so similar to that of  $D_2$  that they will give nearly the same results by this method (see below). The method differs from that of Schleiermacher in that the absolute thermal conductivity of the gas does not have to be considered, but rather the variation of this quantity with the temperature. It may be observed that the temperature variation of the accommodation coefficient should also be taken into account, but since the effect is quite small, the simplified treatment employed above is found to be legitimate.

#### Calibration

The method was tested and the wire calibrated with water containing known amounts, from 3% to nearly 100% of diplogen. These samples were prepared by Dr P. Harteck in the Cavendish Laboratory, and we are most grateful to him for his kindness in supplying them. The water was decomposed on a tungsten wire at  $1000^\circ\text{C}$  (see W, fig 2). It was necessary to ensure complete decomposition of the sample so that there might not be any difference between the D content of the gas and the water from which it was obtained. Tungsten possesses several advantages as a reducing agent, for complete outgassing at  $2000^\circ\text{C}$  eliminates the possibility of any contamination of the recovered gas and the tungstic oxide formed is volatile and distils off so that a fresh metal surface is always being presented to the vapour, while with care it is possible

\* For the exact formula, see A. Farkas, 'Z. phys. Chem.', B, vol. 22, p. 344 (1933).

to obtain 30 c.c. of gas at N.T.P. from a wire 0.2 mm. in thickness and 20 cm long before it burns through. The hydrogen produced was then removed with the aid of a Topley pump (T) and collected in a storage vessel ( $D_2$ ) from which it could be admitted to the measuring cell.

The cell (C) possessed a volume of about 10 c.c., the gas was admitted through the mercury pump  $P_1$ , fig. 2, and the pressure in the cell was adjusted

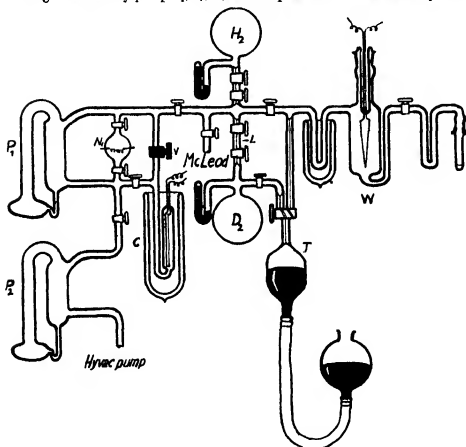


FIG. 2

to the right range with the aid of a carefully ground needle valve (V). A volume of about 0.002 c.c. of gas at N.T.P. is required for a single measurement.\*

\* It is seen from fig. 2 that the sample of gas did not fill only the cell but the dead space of the pump as well. Naturally it would be possible to reduce this volume and to carry out the measurements at a still lower pressure than 0.04 mm. Hg using a thinner wire if it were found desirable to reduce still further the amount of gas required for a single measurement.

which may, with practice, be made in about 2 minutes. This small amount of gas was measured by means of a capillary (L) having a volume of 0.1–0.2 c.c.

The actual readings of the resistances of the wire in the conductivity cell corresponding to the temperatures  $T_2$  and  $T'_2$ , fig. 1, were found to be —

For normal hydrogen—104, 55 ohms

For heavy hydrogen—110, 44 ohms

Thereafter these figures will be referred to as the resistance values of the respective hydrogen-deutrogen sample. In fig. 3 the calibration curve obtained in this way is shown.

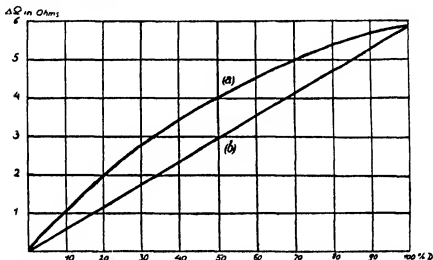


Fig. 3.—Calibration curve. (a)  $H_2 + D_2 + HD$  in equilibrium, (b)  $H_2 + D_2$ .

The straight line shows the relationship obtained between the resistance of the wire and the D content for mixtures containing only  $D_2$  and  $H_2$  and none of the mixed HD molecules, such mixtures having been prepared from normal  $H_2$  and pure  $D_2$ . The curved line shows the resistance obtained when  $H_2$ ,  $D_2$ , and HD are all present in thermodynamical equilibrium (see below), these mixtures having been prepared either by passing the gas over a catalyst or by the complete decomposition of a sample of water of known D content. The difference between the curves is due to the fact to which attention has already been called, namely that the HD and  $D_2$  are so similar in their thermal conductivity variation with temperature that they give nearly the same effect in the measuring vessel, and thus the total HD +  $D_2$  content increases faster with small D contents than with higher ones.

*The Equilibrium  $H_2 + D_2 \rightleftharpoons 2HD$* 

Since the formation of the equilibrium mixture from a pure  $H_2$  and  $D_2$  mixture results in a marked increase in the resistance value of the hydrogen under examination, the technique described above permits the percentages of all three molecules in the mixture to be determined in any given experiment. The formation of the equilibrium mixture containing HD molecules from a 50%  $H_2$  and  $D_2$  mixture results in an increase of the resistance value by about 1 ohm. To determine the quantities of all three gases, it is only necessary after the first measurement of the resistance value, to cause the equilibrium mixture to be formed, return the gas to the cell, and re determine its resistance value, the difference between these quantities will then give the amount of HD formed.

The theoretical equilibrium between the three molecules  $H_2$ ,  $D_2$ , and HD has been discussed by Urey and Rittenberg\*. The equilibrium constant

$K = \frac{[HD]^2}{[H_2][D_2]}$  in the range 200° K to 800° K is given by the expression

$$- \ln K = \frac{\Delta E_0}{RT} - \frac{1}{2} \ln \frac{M_{HD}^2}{M_{H_2} M_{D_2}} - \ln \frac{J_{HD}^2}{J_{H_2} J_{D_2}} - \ln^4$$

where  $J$  denotes the respective moments of inertia,  $M$  the masses and  $\Delta E_0$  the heat of the reaction.  $\Delta E_0$  is about 180 cal, and is the difference in the zero point energy ( $\epsilon$ ) of the molecules taking part in the reaction,  $\epsilon$ ,

$$\Delta E = 2\epsilon_{HD} - (\epsilon_{H_2} + \epsilon_{D_2}),$$

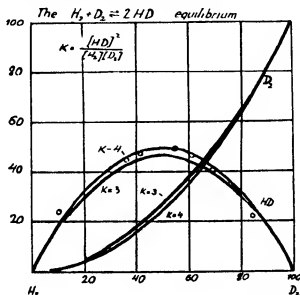
since  $\Delta E_0$  is so very small,  $K$  is practically independent of the temperature.

Experimental examination of the theoretical equilibrium constant in the following manner was carried out†. The increase in the resistance value of a 50%  $H_2$ -50%  $D_2$  mixture when equilibrium has been attained has been assumed to be proportional to the theoretical value, and the single resistance value so obtained is then used to calibrate the wire in terms of the amounts of HD formed in all the other mixtures. The maximum increase in the resistance value results from the formation of HD from a 50%  $H_2$ , 50%  $D_2$  mixture, and in our measuring cell is 1.07 ohms, so that approximately 1% HD will give

\* 'J Chem Phys,' vol 1, p 137 (1933)

† Since the first publication of our results ('Nature,' vol 132, p 894 (1933)), Rittenberg, Bleakney, and Urey ('J Chem Phys,' vol 2, p 48 (1934)) have published a note showing that they have succeeded in measuring the equilibrium constant with a higher accuracy by mass spectroscopic methods.

an increase of 0.02 ohms. Fig. 4 shows that with this procedure the amounts of HD obtained when equilibrium from various mixtures of  $H_2$  and  $D_2$  is obtained lie with remarkable accuracy on the theoretical curve. In this particular experiment it was desired to test the applicability of the method of measurement, rather than to determine the absolute value\* of the equilibrium constant, but attention may be called to the fact that these experiments show that there is no appreciable alteration in the equilibrium constant with temperature, for when an equilibrium mixture prepared at 20° C (by the decomposition



of heavy water by sodium) is heated to 600° C. on a nickel wire no appreciable alteration in the resistance value is observed.

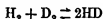
It is found that the formation of the equilibrium mixture at 600° C. on a nickel wire occurs in a few seconds. Thus to determine the amount of HD in a sample, its resistance value having been determined, it was pumped from the measuring cell into a vessel containing a hot nickel wire, Ni, fig. 2, left in contact with the wire for a few seconds, pumped back to the measuring cell and its resistance value again determined. With this procedure, it is important that the nickel wire should not retain or evolve any hydrogen, so that the

\* In some experiments to be published in detail later (cf. 'Nature,' *loc. cit.*) on the rate at which this equilibrium is attained the absolute value of  $K$  need not be known, for only the relative change is under consideration.

amount of the gas used in the two measurements is the same. The wire must be maintained scrupulously clean, and have been perfectly outgassed, for every trace of foreign gas present in such minute amounts of material would cause a large error in the measurement.

More detailed measurements of the formation of HD from  $H_2$  and  $D_2$  on nickel and other metals will be carried out shortly, but it is already possible to propose a mechanism for this process. The hydrogen molecules are adsorbed and dissociated into atoms on the surface of the metal, and naturally on desorption combine in the adsorption layer to give  $H_2$ ,  $D_2$ , and HD molecules whose relative amounts are determined by the value of this equilibrium constant at the temperature of the catalyst. This mechanism is identical with the so-called high temperature mechanism of the catalytic conversion of para hydrogen.\*

We have also attempted to bring about the reaction



at low temperatures, by adsorbing the gases on charcoal but with negative results. 0.01 cc of a 50%  $H_2$  and 50%  $D_2$  mixture were adsorbed for 15 hours on 20 mg charcoal at 77.3° K, and on desorption no formation of HD could be detected, though in theory at this temperature the equilibrium concentration is about 40%. This negative result shows rather beautifully that adsorption or desorption on charcoal causes no rupture of the molecules for if such did actually occur an exchange of D and H atoms would take place in the surface layer resulting in the formation of HD. The work of Bonhoeffer, Farkas and Rummel on the mechanism of the heterogeneous conversion of para- to ortho-hydrogen would also lead us to anticipate such a result, for the latter reaction is unimolecular in the adsorbed phase and involves no exchange mechanism.†

Since it is apparent that the equilibrium at low temperatures is not attainable by means of a catalyst, it will be necessary, if measurements in this region are to be made, to obtain the diplogen-hydrogen-mixture *in statu nascendi* either photochemically or by a suitably chosen chemical reaction. On recombination the equilibrium mixture of  $H_2$ ,  $D_2$ , and HD will be obtained. Such measure-

\* Cf. Bonhoeffer, Farkas, and Rummel, 'Z. phys. Chem.,' B, vol. 21, p. 225 (1933).

† The establishment of the ortho- and para-hydrogen equilibrium on charcoal results from the inhomogeneous magnetic field set up by unsaturated para-magnetic carbon atoms in the surface allowing a reversal of the nuclear spin to take place. On the above-mentioned charcoal the ortho-para-hydrogen equilibrium was established within a few minutes at 78° K.

ments would, as Urey and Rittenberg (*loc cit*) have already pointed out, have particular interest since they should give direct information on the nuclear spin and the statistics of the molecules under consideration

### *Separation by Diffusion*

Certain irregularities which we had observed in measurements of the concentrations of various mixtures led us to examine the possible separation of the isotopes occurring as a result of their different molecular velocities. Such separation might be expected to occur when the mixed gases stream at low pressures through such narrow openings as partially closed taps or valves, and was, in fact, particularly marked where the measuring cell had initially been filled with too great a charge of hydrogen and its pressure later adjusted to the correct value by pumping off the excess gas through a very fine regulating valve. A concentration of the heavy isotope takes place in the cell while the dihydrogen content of the gas pumped off is naturally less. The magnitude of the effect may be calculated from Rayleigh's formula for fractional distillation which is valid for the range of pressure where the mean free path of gases effusing from a narrow opening is much greater than the diameter of the nozzle

$$\frac{(H_0)}{(D_0)} \cdot \frac{(H)}{(D)} = \left(\frac{p_1}{p_0}\right)^{s-1},$$

( $H_0$ ), ( $D_0$ ), ( $H$ ) and ( $D$ ) denoting the initial and the final concentration of the hydrogen,  $s$  being the ratio of the molecular velocities,  $\pm e$ , in this case  $\sqrt{2}$ ,  $p_0$  the initial pressure and  $p$  the final pressure equal to 0.04 mm. Hg,  $\pm e$ , the same pressure which was maintained in the conductivity cell in all measurements

Table I shows that the theory accounts well for the results obtained

Table I

$p_0/p$	% D observed	% D calculated
1	47.5	—
1.5	50.7	50.7
2	53.0	53.0
3.25	56.0	57.8

The small discrepancy in the last experiment is probably due to the fact that the pressure at which the experiment was carried out (about 0.13 mm Hg) was too high for the nozzle of the valve, and the conditions of ideal effusion were no longer maintained.

A similar effect is also to be expected when  $H_2$ ,  $D_2$  and HD are all present in their equilibrium concentrations, but now the theoretical treatment of the problem is more involved, and the estimation of the concentrations is technically more difficult since the equilibrium is somewhat modified through the separation that takes place. The effect is, however, revealed in the alteration in the resistance of the wire on pumping off the gas, and naturally such an effect is not to be anticipated when pure hydrogen or a high percentage diplogen concentrate are in the cell. No such alteration is actually observed under these conditions, and this may be used as a convenient test for the purity of the diplogen. In all the later measurements careful control of the dimensions of the apparatus in all parts where the flow of gas was at all restricted, and the admission of only just the right amount of gas to the cell ensured that all such variations in the measurements of the concentrations were eliminated.

No preferential adsorption of the diplogen occurs on exposure of the two hydrogen isotopes to charcoal. In an experiment in which 96% of a 30% hydrogen-diplogen mixture was adsorbed on charcoal at 78° K, the gas remaining above the charcoal had the same H/D ratio as the original mixture. Enrichment of the D in the mixture residue occurs only if the gas above the charcoal is pumped off, and is due to the different diffusion velocities of the three molecular species present.

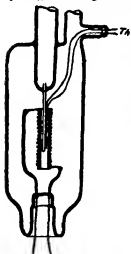


FIG. 5—Th = thermo-element.

#### *The Diffusion of the Isotopes through Palladium*

The diffusion of the two hydrogen isotopes from mixtures containing 40% to 50% of D through palladium at different temperatures has been studied.

The gas at a pressure of about 5 mm. was allowed to diffuse through a thin hot palladium tube at constant temperature, fig. 5, as measured by a thermo-element, into a vacuum from which it could be pumped off into the measuring vessel and its composition determined. From time to time measurements were also made of the concentration of the gas remaining behind. It was



found that the gas which had diffused through the palladium always had a lower D content than the original mixture, but that the difference in the D contents of the original gas and that of the gas diffused through the palladium decreased as the temperature of the palladium tube was increased. Further the H/D ratio of the diffused gas did not depend on whether the initial mixture was composed only of  $H_2$  and  $D_2$  or of mixtures of  $H_2$ ,  $D_2$ , and HD of the same H/D ratio\*. This would naturally be expected, since it is sure that inside the metal all these molecules are fully dissociated. Accordingly, the gas diffusing through, and also after a time the initial sample, contains the equilibrium mixture of the three molecules. In fig. 6 the ratio

$$\alpha = \frac{\left(\frac{H}{D}\right)_{\text{diffused}}}{\left(\frac{H}{D}\right)_{\text{original}}}$$

is shown as a function of the temperature

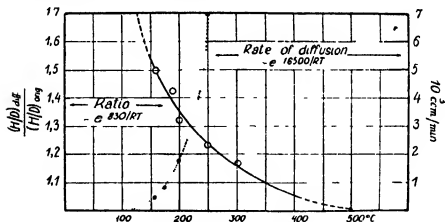


FIG. 6—Pressure 5 mm Hg

The figure shows that at higher temperatures the curve becomes asymptotic approaching unity as a limit, indicating that then the H and D are diffusing through the palladium with equal velocities. The ratio gives directly the relative velocities of the two forms.

\* For diffusion through a thin quartz tube, this would not necessarily be true, since in this experiment molecules might diffuse through without previous dissociation.

The dependence of the diffusion ( $v$ ) of H and D through palladium with temperature can be expressed by the relation

$$v = v_H + v_D = C e^{-A/RT},$$

where  $A$  is the so-called activation energy of diffusion.

The dotted curve in fig 6 shows these amounts plotted against the temperature, for which  $A$  has the value 16,500 cal. The amounts of H and D which diffuse through palladium may, also to a first approximation, be expressed by the two equations

$$v_H = v(H)_{diff} = C_H (H)_{orig} e^{-A_H/RT},$$

$$v_D = v(D)_{diff} = C_D (D)_{orig} e^{-A_D/RT}$$

From these two expressions the magnitude

$$\alpha = e^{(A_D - A_H)/RT}$$

is obtained and is plotted in fig 6 (provided that the constants  $C_H$  and  $C_D$  in both expressions are equal, see below). From the values shown in the figures  $A_D - A_H = 830$  cal., which value is then the excess of the activation energy for the diffusion of D over that of H and is, in fact, about 5% of the whole (The value  $A = 16,500$  cal. is the average of  $A_H$  and  $A_D$ ).

As a result of the difference in the velocities of diffusion of H and D the hydrogen in the original vessel from which diffusion is taking place gets gradually richer in the heavy isotope. It is easily seen that the amount of concentration is again given by the Rayleigh formula.

Fig 6 shows that at temperatures higher than  $300^\circ\text{C}$   $\alpha$  is very nearly equal to unity, and thus at these temperatures no marked separation is to be anticipated. It is for this reason that attempts to use the diffusion of hydrogen through palladium as a means of concentrating the isotope have failed. At lower temperatures, when the separation factor is relatively great, the amount of gas passing through is unfortunately so small that any separation by simple methods such as passing hydrogen at atmospheric pressure through a tube of ordinary dimension is quite impracticable.

The difference between the two diffusion velocities may be considered to arise in the following ways

Fig 7 shows the energy levels of the two isotopes during the sorption process. The difference in energy of the  $\text{H}_2$  and  $\text{D}_2$  gases arises from the difference of their zero-point energies. The level for  $\text{HD}$  lies fairly exactly midway between that for  $\text{H}_2$  and  $\text{D}_2$ . In the sorbed state the gases are dissociated

into atoms and corresponding to the different sorption complexes Pd-H and Pd-D, have still different zero-point energies

In the stationary state the velocity of diffusion must be determined by the velocity of sorption, as the same amount of gas must always be taken up as is being desorbed. Thus it is clear that the velocities of sorption of the two hydrogen isotopes must actually be different because of the different activation energies, which have been shown to differ by about 830 cals, i.e., about the same as the difference in the zero point energy of  $\frac{1}{2}\text{H}_2$  and  $\frac{1}{2}\text{D}_2$ , i.e., 890 cals. None the less there must certainly be a difference of the same order of magnitude in the heats of adsorption of the two molecules, resulting in a greater solubility or sorption of the heavier than the lighter isotope.

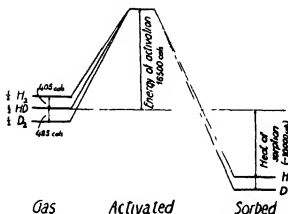


FIG. 7

This consideration is merely schematic with a very simplified model, for it is assumed that the rate of diffusion is not determined by the migration of the hydrogen through the lattice (this is true for an infinitely thin palladium layer) or that, at least, the rates of migration are equal for both hydrogen isotopes.

In this connection also, it can be reasonably suggested that the electrolytic separation owes its efficacy to a similar effect, and it is remarkable that if the factor  $\alpha = e^{-(A_D - A_H)/RT}$  is extrapolated to room temperature a separation factor of five results, in agreement with the results first obtained by Lewis and Macdonald\*. It should be observed that the difference in the energies of activation will be nearly independent of the metal surface, for the zero point energies of the various metal-hydrogen complexes will vary only slightly from

\* J. Chem. Phys., vol. 1, p. 345 (1933)

metal to metal, thus the separation factor on different metals by electrolysis will be nearly independent of the metal surface, a conclusion in agreement with the experiments of Bell and Wolfenden \* †

We are very much indebted to Professor E K Rideal, FRS, both for discussions and for his interest in our work, to Dr O H Wansbrough-Jones for translation of our manuscript and to the Central British Fund for German Jewry for a financial grant

### *Summary*

(1) A method based on the different specific heats of the hydrogen isotopes is described permitting a determination of the concentration of the heavy hydrogen isotope in a mixture with an accuracy of 0.2% in a sample of 0.002 c.c. N.T.P. of gas

(2) It is shown that in a mixture of hydrogen and diplogen the equilibrium



is readily established at the surface of a hot nickel wire. The equilibrium constant is, in agreement with theory, about 4 and nearly independent of temperature

(3) This establishment of the equilibrium is used to estimate the ratios of the molecules  $H_2$ ,  $D_2$ , and  $HD$  in a given sample of hydrogen

(4) By pumping hydrogen-diplogen mixtures at low pressures through a fine nozzle a slight separation occurs due to the different molecular velocities of the hydrogen isotopes

(5) If a hydrogen-diplogen mixture diffuses through a palladium tube the  $H/D$  ratio of the diffusing gas is the greater the lower the temperature of the tube. Above  $300^\circ C$  the  $H/D$  ratio of the diffusing gas is equal to that of the original gas. This may be explained by assuming that the energy of activation for the diffusion of diplogen is greater than that for hydrogen on account of the difference in their zero point energies

\* 'Nature,' vol 133, p 26 (1934)

† Topley and Eyring, 'Nature,' vol 131, p 292 (1934), 'J Amer Chem' vol 55, p 5058 (1933)

## *Experiments on Heavy Hydrogen 11—The Ortho-para Conversion*

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(Communicated by Eric K Rideal, F R S—Received March 2, 1934)

Ortho- and para-modifications should exist in all diatomic molecules of identical atoms with nuclear spin, since the law of prohibition of the ortho-para intercombination is generally applicable\*. The rotational states consist alternately of ortho- and para-levels the ortho-levels being those which possess on the average the higher quantum weight and represent the states with the symmetrical nuclear-spin eigenfunctions, whereas the para-levels correspond to the antisymmetrical nuclear-spin function.

The rotational quantum numbers possessed by the ortho- and para-levels depend on the statistics valid for the nuclei and on the symmetry of the electronic eigenfunctions. For even electronic eigenfunction (for hydrogen and diplogen the ground states of which are  $^1\Sigma_g$ ) if the Fermi-Dirac statistics are valid the ortho-modifications are in the rotational states with odd quantum numbers and the para states in the even ones, whilst the Bose-Einstein statistics lead one to expect the reverse to be true.

The existence of the two sets in the molecule becomes apparent on studying the specific heats,† the spectrum,‡ and the equilibrium of the two modifications.§ With ordinary hydrogen these three effects were observed in that order, for diplogen it has already been shown that alternating intensities occur in the emission spectrum|| and in this paper we shall substantiate the differences in the rotational specific heat and examine the ortho-para equilibrium.

The ratio of the concentration of the ortho- and para-modification in equilibrium with each other may be derived in the following way¶ let  $t$  be the

\* Cf e.g. L. Farkas, 'Ergbn exact Naturw.', vol 12, p 163 (1933)

† Dennison, 'Proc Roy Soc.' A, vol 115, p 483 (1927), Eucken, 'SitzBer Preuss Akad. Wiss. Berlin,' No 141, p 141 (1912)

‡ Heisenberg, 'Z. Physik,' vol 38, p 41 (1926), vol 41, p. 239 (1927), Hund, 'Z. Physik,' vol 42, p 93 (1927)

§ Bonhoeffer and Harteck, 'Z. phys. Chem.' B, vol 4, p 113 (1929), Eucken and Hiller, 'Z. phys. Chem.' B, vol 4, p 142 (1929)

|| Lewis and Ashley, 'Phys. Rev.', vol 43, p 837 (1933)

¶ Weizel, "Bandenspectren," 'Wien-Harms Handbuch der Exp. Physik,' (1933)

nuclear spin of the atom then the resulting spin  $T$  of the molecule may have  $2t + 1$  different values

$$T = 2t, \quad \underline{2t - 1}, \quad 2t - 2, \quad \underline{2t - 3}, \quad \dots, \quad 0$$

the values underlined corresponding to the antisymmetrical spin function, i.e., to the para states. As each resulting nuclear spin  $T$  is  $2T + 1$  fold degenerated, the normal or high temperature ortho-para ratio is given by the ratio of the total multiplicity ( $g$ ) of the ortho states to that of the para states, viz.,

$$\frac{g_{\text{ortho}}}{g_{\text{para}}} = \frac{\sum_{n=\text{even}} 2(2t - n) + 1}{\sum_{n=\text{odd}} 2(2t - n) + 1} = \frac{(t + 1)(2t + 1)}{t(2t + 1)} = \frac{t + 1}{t}$$

The distribution of the molecules in different rotational levels and thus the dependence on temperature of the ortho-para equilibrium is governed by the Boltzmann law

$$N_j = N_0 g(2j + 1) e^{-E_j/kT},$$

$N_j$  denoting the fraction of the molecule of the total number ( $N_0$ ) in the rotational level  $j$  with the energy

$$E_j = \frac{j(j + 1) \hbar^2}{8\pi^2 J}$$

( $J$  = moment of inertia),  $g$  is the nuclear statistical weight, being  $(t + 1)(2t + 1)$  for the ortho states, and  $t(2t + 1)$  for the para states.

The fraction of the molecules in the para and ortho states is given by

$$b_{\text{para}} = \frac{\sum_{\text{para}} g_{\text{para}}(2j + 1) e^{-E_j/kT}}{\sum_{\text{ortho}} g_{\text{ortho}}(2j + 1) e^{-E_j/kT} + \sum_{\text{para}} g_{\text{para}}(2j + 1) e^{-E_j/kT}} = \frac{Q_{\text{para}}}{Q_{\text{ortho}} + Q_{\text{para}}}$$

$$b_{\text{ortho}} = \frac{Q_{\text{ortho}}}{Q_{\text{ortho}} + Q_{\text{para}}}$$

The index ortho or para under  $\Sigma$  denotes that the summation must be extended over the ortho or para states.

Experiments were carried out in order to demonstrate the ortho-para conversion in diplogen and to determine from the equilibrium ratio the spin and statistics of the diplogen nucleus.

The experiments were planned on the assumption that diplogen would behave in a manner analogous to hydrogen, that is to say, that the attainment of the equilibrium ratio which, on account of the impossibility of the ortho-para intercombination taking place in pure hydrogen, is extremely slow, could

be accelerated by suitable catalysts and that the gas removed from the catalyst, being stable under ordinary conditions, retains the ortho-para concentration which is in equilibrium at the temperature of the catalyst

As catalysts for the rapid production of the ortho para diplogen equilibrium mixture an active charcoal, which effected a very rapid ortho para conversion in hydrogen even at the temperature of liquid hydrogen, was employed. The sample of diplogen was of about 90% purity. It is evident that these experiments could be performed more readily with pure diplogen for the 90% diplogen contains the following five constituents: ortho hydrogen, para-hydrogen, HD molecules, ortho-diplogen, and para-diplogen. It was, however, found possible to determine the concentration of each constituent and thus overcome this difficulty.

For the determination of the different forms of hydrogen and diplogen the micro method of A. Farkas,\* which gives measurements of high precision with amounts of gas as small as 0.002 cc at N.T.P., was employed. This method was applicable since a difference in the rotational specific heats of ortho- and para-diplogen was to be expected similar to that found in ortho- and para-hydrogen (cf fig. 2). The actual measurements were carried out in two thermal conductivity cells having wires of different length and thickness extended down the axis of each cell.

The calibration readings† were as follows:—

Table I

	Cell I	Cell II
	ohms	ohms
Normal hydrogen	105.23	128.39
Para hydrogen	106.45	131.44
Normal diplogen (90% D)	111.23	136.73

For hydrogen it was shown that for these cells the excess concentration of para-hydrogen relative to normal hydrogen was proportional to the corresponding differences in the values found for the resistance, according to the formula

$$(p - 25) = \text{prop} (R_p - R_{25}),$$

where  $p$  denotes the para-hydrogen concentration in percentage of the sample,  $R_p$  its resistance value and  $R_{25}$  the resistance value of normal or 25% para-hydrogen.

\* *Z. phys. Chem.*, B, vol. 22, p. 344 (1933).

† A. Farkas and L. Farkas, *Nature*, vol. 132, p. 894 (1933).

To determine the exact equilibrium between ortho- and para-diplogen three series of experiments were carried out at the temperatures 20 °K, \* 53° K and 78° K. The first run was carried out at 20 °K. The catalyst consisting of about 0.2 gram of active charcoal, enclosed in a thin glass tube of a few cubic centimetres capacity, absorbed at 20 °K about 50 c.c. of diplogen, but the pressure over the charcoal was not higher than a few tenths of a millimetre of mercury. The first samples of gas were extracted from the charcoal tube after the diplogen had been in contact with the catalyst for some minutes, and its resistance value was found to be (cell I) 111.58 ohms compared with 111.23 ohms for the original gas.

A change in the resistance value may be due to four different causes —

- (1) Through separation of the hydrogen isotopes on pumping off the samples owing to the higher diffusibility of the lighter hydrogen, Part I
- (2) Through the possible readjustment of the  $\text{H}_2 + \text{D}_2 \rightleftharpoons 2\text{HD}$  equilibrium
- (3) Through the ortho-para conversion of the ordinary hydrogen present in the original sample or formed by reaction (2)
- (4) Through the ortho-para diplogen conversion

From the calibration experiments, it is evident that a separation of the hydrogen isotope in pumping off the samples cannot cause a rise in the resistance value but must lower it owing to the lower diplogen content of the gas pumped off.

Previous experiments, made on charcoal indicated that the readjustment of the  $\text{H}_2 + \text{D}_2 \rightleftharpoons 2\text{HD}$  equilibrium did not occur at the temperature of liquid nitrogen and, therefore, it is very improbable that the change in the resistance value is due to such an effect at the temperature of liquid hydrogen, but if (2) does not occur there is very little probability that the effect found is due to the ortho-para conversion of ordinary hydrogen since the original gas (90% D) could not contain more than 1%  $\text{H}_2$ , and such a small amount (even if somewhat concentrated by the pumping) would not cause the observed change in the resistance value when converted to para-hydrogen.

This single measurement, therefore, makes it very probable that we are dealing with a true para-ortho diplogen conversion. This was confirmed by the following experiment.

\* The liquid hydrogen was kept in a liquid air cooled Dewar container of 350 c.c. By daily renewal of the liquid air the liquid hydrogen could be preserved for 4 days.



The gas pumped off from the charcoal cooled by liquid hydrogen was re-adsorbed on charcoal at 78° K and then completely re-evaporated, this sample showed a resistance value of 110.75 ohms, i.e., 0.83 ohms lower than the sample taken from the charcoal at 20.4° K. In this case it is evident that the processes (1) and (2) could not occur, the actual fall in the resistance value was, therefore, due to the adjustment of the ortho-diplogen para-diplogen equilibrium from 20.4° K to 78° K.

We cannot attribute this change in the resistance value to the hydrogen content of the sample, for so great a change could only be caused by the ortho-para conversion of pure hydrogen. Finally, this sample of gas was brought into contact with a hot nickel wire and the resistance value obtained after this operation was 110.25 ohms. This further fall was to be anticipated since the excess of HD contained in the sample pumped off the charcoal cooled by liquid hydrogen was transformed according to the equation



From this final resistance value we calculate the D content to be about 67%\* and from the difference 110.75-110.25 after taking into account a small change due to the readjustment of the ortho-para-equilibrium of diplogen (see below) we find that about 12% HD has disappeared, i.e., when the original gas had a composition of

$$90\% \text{ D} = 81\% \text{ D}_2 + 18\% \text{ HD} + 1\% \text{ H}_2,$$

the gas pumped off from the charcoal cooled by liquid hydrogen contained

$$67\% \text{ D} = 39\% \text{ D}_2 + 56\% \text{ HD} + 5\% \text{ H}_2$$

the equilibrium mixture being

$$45\% \text{ D}_2 + 44\% \text{ HD} + 11\% \text{ H}_2$$

This procedure was repeated several times and always led to practically the same result and conclusion. Since, even by continuous pumping, we failed to get off the charcoal cooled in liquid hydrogen a gas containing more than 70% D (the whole amount pumped off in 10 hours was not more than 1/10 of that originally adsorbed) the liquid hydrogen container was suddenly removed and the greatest part of the adsorbed gas was recovered. In the earlier experiments of Bonhoeffer and Harteck† with para-hydrogen, it was found

\* This separation compared with 90% D of the original gas is due to a slightly preferential adsorption which does not occur at higher temperatures.

† 'Z. phys. Chem.,' B, vol. 4, p. 113 (1929)

that when para-hydrogen is suddenly liberated in this manner the ortho-para hydrogen equilibrium is not readjusted to the actual temperature of the charcoal as the desorption takes place with a greater velocity than the re-conversion

A sample of this gas gave a resistance value of 112.61 ohms which was reduced by contact with a hot nickel wire to 111.25 ohms, *i.e.*, practically to the value of the original sample. The corresponding readings in cell II were 138.93 ohms and 136.73 ohms respectively.

As no separation whatever could occur the whole change was entirely due to the ortho-para diplogen conversion.

To determine the temperature dependence of the ortho-para diplogen equilibrium further series of measurements were made at 53° K and 78° K. In Table II the changes are recorded in the resistance value for the samples relative to the normal diplogen and their ratios.

Table II

Temperature, °K	Cell I *		Cell II	
	Change	Ratio	Change	Ratio
20.4	1.40	9.2	2.20	8.8
53	0.51	3.4	0.86	3.4
78	0.15	1	0.25	1

\* More weight is to be assigned to the values from cell I than to those from cell II, since they include a greater number of measurements.

Since previous experiments with hydrogen had shown that the changes in the resistance value were proportional to the changes in the para-hydrogen content of the gas, an attempt was made to estimate from these readings the actual concentration of the ortho and para component in the following way.

Fig. 1 shows the dependence of the concentration of ortho-diplogen on the temperature adopting the Dirac-Fermi or Bose-Einstein statistics and for nuclear spins of  $\frac{1}{2}$  and 1 (higher values of  $i$  are most improbable). Table III contains the ratios of the excess concentration of ortho- and para-diplogen at 20.4° K, 53° K, and 78° K.

If we compare the relative changes in the concentration with the ratio of the changes in the resistance value we can certainly exclude the Dirac-Fermi statistics. If the Bose-Einstein statistics are applicable to diplogen then the lowest rotational level with the quantum number 0, and thus all rotational states with even quantum numbers, are ortho states, and have the higher

statistical weight This result is in agreement with the statement of Lewis and Ashley who found that in the emission spectrum of the diplogen molecule the lines in the Q-branches of the Fulcher-bands corresponding to the transitions between rotational levels with even quantum numbers are more intense Our ratios agree better with the spin  $t = 1$  which is the value to be expected from the analogy with other elements, *i.e.*, those with even atomic weight having integer nuclear spins \*

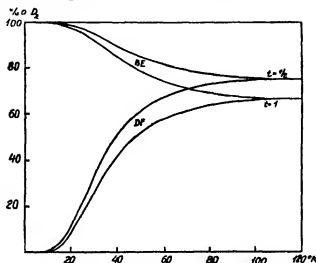


FIG 1—BE calculated according to the Bose Einstein statistics, DF according to the Dirac Fermi statistics

Table III

Temperature, °K.	Dirac Fermi statistics		Bose Einstein statistics	
	$t = \frac{1}{2}$	$t = 1$	$t = \frac{1}{2}$	$t = 1$
20.4	31.2	14.7	8.4	9.5
53	3.8	3.2	1.1	3.2
78	1	1	1	1

For the nuclear spin 1 the lowest rotational state with the quantum number 0 has the following constitution The spins of the nuclei are either parallel

\* It may be added that the measurements cannot exclude definitely a nuclear spin of  $\frac{1}{2}$  as the relative changes in the concentrations (in this substance) are very similar to those for  $t = 1$  The same difficulty appears in the spectroscopic measurement of the alternating intensity A conclusive determination of the nuclear spin can only be made by the absolute measurement of the specific heats as was carried out by Eucken for hydrogen

or antiparallel having the resultant spin  $T = 2$  or  $0$ , respectively. The state with  $T = 2$  is  $2T + 1 = 5$ -fold whereas the state  $T = 0$  is single. The latter has the lowest energy when this 6-fold degenerated state is split up at extremely low temperatures for example in the crystal. Transition between these two sets would also involve a change in the multiplicity of the nuclear spins similar to the para-ortho hydrogen transition, we should therefore expect a prohibition for this transition.

If we compare the dependence on temperature of the ortho-diplogen concentration with hydrogen we note a remarkable difference owing to the different statistics while in hydrogen the ortho-hydrogen, i. e., the component present in excess at high temperatures, disappears on lowering the temperature, in diplogen the ortho modification is the only stable form at low temperatures.

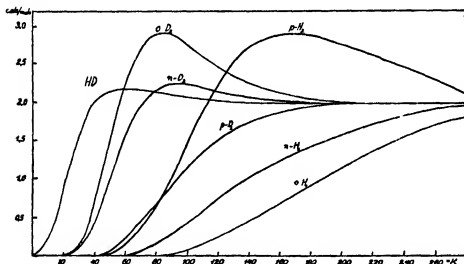


FIG. 2

In fig. 2 the rotational specific heats curves are shown for normal diplogen ( $n\text{-D}_2$ ), ortho-diplogen ( $o\text{-D}_2$ ), para-diplogen ( $p\text{-D}_2$ ), normal hydrogen ( $n\text{-H}_2$ ), para-hydrogen ( $p\text{-H}_2$ ), ortho-hydrogen ( $o\text{-H}_2$ ) and for the mixed molecule gas HD. As normal diplogen and normal hydrogen are designated those mixtures of ortho- and para-molecules which correspond to the distribution at high temperatures, i. e.,

$$n\text{-D}_2 = \frac{2}{3} o\text{-D}_2 + \frac{1}{3} p\text{-D}_2,$$

$$n\text{-H}_2 = \frac{2}{3} o\text{-H}_2 + \frac{1}{3} p\text{-H}_2.$$

The specific heat curves were calculated on the basis of the Dennison formulæ (loc cit)

$$C_{\text{rot}}^{\text{ortho-D}_2} = R\sigma^2 \frac{d^2 \ln Q_{\text{para}}}{d\sigma^2},$$

$$C_{\text{rot}}^{\text{ortho-H}_2} = R\sigma^2 \frac{d^2 \ln Q_{\text{ortho}}}{d\sigma^2},$$

$$C_{\text{rot}}^{\text{ortho-D}_2} = \frac{2}{3} C_{\text{rot}}^{\text{ortho-D}_2} + \frac{1}{3} C_{\text{rot}}^{\text{ortho-H}_2} \quad \text{with } \sigma = \frac{\sigma_{\text{H}_2}}{2} = \frac{85}{2T},$$

on account of the double moment of inertia of  $\text{D}_2$  compared with  $\text{H}_2$ . For HD the method of computation was based upon the analysis of Gregory.\*

A comparison has been made between the reaction velocity of the ortho-para conversion of diplogen in two characteristic examples with the corresponding reaction of hydrogen, firstly the heterogeneous ortho-diplogen formation at  $20.4^\circ \text{K}$ . and secondly the re-conversion of ortho-diplogen at room temperatures catalysed by gaseous oxygen.

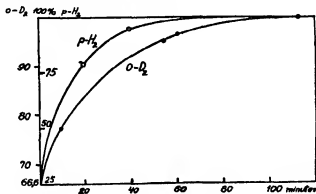


FIG. 3.

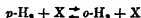
Since, as we have seen, the formation of ortho-diplogen is very rapid on charcoal another catalyst was employed consisting of a glass tube which, owing to some impurities, catalysed the para-ortho-diplogen conversion with a conveniently measurable speed. On this catalyst the diplogen conversion proceeded slightly more slowly than the hydrogen conversion as shown in fig. 3, namely, with a half life time† of about 19 and 11 minutes respectively.

\* 'Z Physik,' vol. 78, p. 791 (1932).

† As was shown for hydrogen this type of reaction proceeds according to the formula  $p_t - p_\infty = (p_0 - p_\infty)e^{-kt}$ ,  $p_0$ ,  $p_t$ , and  $p_\infty$  denoting the para hydrogen concentration at a time  $t=0$ ,  $t$  and  $t=\infty$ . Then the reaction rate is given by the half life  $\tau = \ln 2/k$ .

The mechanism of the catalytic ortho-para conversion of diplogen is most probably the same as for hydrogen,\*† i.e., we have again two independent mechanisms one of them acting at low temperatures and the other at high temperatures, one is connected with the normal or van der Waals' adsorption and the second with the chemi- or atomic adsorption. The low temperature mechanism (e.g., catalysis by charcoal) involves only one molecule and the twisting of the nuclear spin from the para position to the ortho position is effected by the magnetic forces of the catalyst. The probability of this twisting is nearly independent of the temperature and the reaction rate is governed by the amount adsorbed. The high temperature mechanism (e.g., catalysis by metals) consists in atomic adsorption of diplogen (or hydrogen) when the adsorbed molecules are split into atoms and the ortho- and para-molecules re-evaporate in a ratio corresponding to the equilibrium at the actual temperature of the catalyst.

The homogeneous reconversion of ortho-diplogen in the presence of oxygen was next investigated, for it provides us with valuable information on the value of the nuclear magnetic moment of the diplogen. As has been recently shown by L. Farkas and Sachse† paramagnetic molecules or ions convert para-hydrogen to normal hydrogen according to the reaction



The rate of the reaction is given by

$$p_t - p_\infty = (p_0 - p_\infty) e^{-(k_{p \rightarrow o} + k_{o \rightarrow p})[X]t},$$

where  $k_{p \rightarrow o}$  and  $k_{o \rightarrow p}$  are the reaction velocity constants of the respective reactions and  $[X]$  is the concentration of the paramagnetic substance. At temperatures above 200° K  $k_{p \rightarrow o}/k_{o \rightarrow p}$  is 3, because the ratio  $p\text{-H}_2/o\text{-H}_2$  is maintained at a ratio of 1/3. Since the conversion is effected by the perturbation of the inhomogeneous magnetic field of the paramagnetic molecule upon the nuclear magnetic moment it was to be expected that a similar re-conversion would take place with ortho-diplogen as had been observed with para-hydrogen. For the ortho-diplogen conversion above 100° K we have

$$\frac{k_{o \rightarrow p}}{k_{p \rightarrow o}} = \frac{1}{3}$$

since the composition of normal diplogen is 2/3  $o\text{-D}_2$  + 1/3  $p\text{-D}_2$ .

\* Bonhoeffer and A. Farkas, 'Trans. Faraday Soc.', vol. 28, pp. 242, 561 (1932)

† Bonhoeffer, A. Farkas, and Rummel, 'Z. phys. Chem.', B, vol. 21, p. 225 (1933)

‡ 'Sitzber. Preuss. Akad. Wiss. Berlin,' p. 268 (1933), 'Z. phys. Chem.', B, vol. 23, pp. 1, 19 (1933).

The experimental arrangement for investigating the ortho-diplogen re-conversion was very simple. Ortho-diplogen, prepared at 20 ° K on charcoal, was mixed with oxygen and samples of this mixture were taken from time to time for measurement of the ortho-diplogen concentration. The samples passed through a trap, cooled with liquid hydrogen in order to freeze out the oxygen, before reaching the measuring cell. The mixture of ortho-diplogen and oxygen was kept in a gas burette over mercury and its pressure could be adjusted very conveniently by altering the level of the mercury in the burette.

As mean value for the half life time at 10 mm oxygen pressure (293° K) a value of 2200 minutes was obtained corresponding to a velocity constant of  $\frac{\ln 2 \cdot 22 \cdot 4 \cdot 760}{2200 \cdot 10} \cdot \frac{293}{273} = 0.57$  litre/mole minute. The velocity constant is thus 16 times smaller than the velocity constant of the corresponding hydrogen reaction, namely, 9.16 litre/mole minute.

This great difference in the velocity constant can be interpreted on the basis of the theoretical investigations of Wigner\*. According to this author the theoretical collision efficiency of the ortho-para transition from the rotational state  $j = 0$  to the rotational state  $j = 1$  is given by

$$Z_{01} = \frac{8\pi^2 J_{H_2} \mu_p^2 \mu_x^2}{9h^2 k T a^6} e^{-E_0/kT},$$

where  $J_{H_2}$  is the moment of inertia of  $H_2$ ,  $\mu_p$  the magnetic moment of the proton ( $2.505 \cdot 10^{-28}$  CGS units),†  $\mu_x$  the magnetic moment of the colliding paramagnetic molecule and  $a$  the closest distance of approach on collision. The factor  $e^{-E_0/kT}$  is due to the fact that this transition is endothermic. From the value  $Z_{01}$  the collision efficiency can be derived for the whole para-ortho reaction ( $Z_{p \rightarrow o}$ ) which includes all the para-ortho transitions  $j = 0 \rightarrow 1$ ,  $2 \rightarrow 1$ ,  $2 \rightarrow 3$ , etc., and which is connected with the observed velocity constant by the relation

$$Z_{p \rightarrow o} = \frac{(k_{p \rightarrow o} + k_{i \rightarrow p})}{\text{number of collisions per minute}},$$

and is given by

$$Z_{p \rightarrow o} = \frac{8\pi^2 J_{H_2} \mu_p^2 \mu_x^2}{9h^2 k T a^6} \left( \frac{e^{-E_0/kT} + 2e^{-E_1/kT} + 3e^{-E_2/kT} + \dots}{1 + 5e^{-E_0/kT} + 9e^{-E_1/kT} + \dots} \right)$$

\* 'Z. phys. Chem.,' B, vol. 23, p. 28 (1933).

† Estermann, Frisch, and Stern, 'Nature,' vol. 132, p. 169 (1933), Estermann and Stern, 'Z. Physik,' vol. 85, p. 17 (1933), Frisch and Stern, 'Z. Physik,' vol. 85, p. 4 (1933).

If we employ a similar expression for  $Z_{o \rightarrow p}$ , in the case of ortho-diplogen and adopt the same value of  $\alpha$  in both cases we obtain for the ratio

$$\frac{Z_{p \rightarrow o}^{H_2}}{Z_{o \rightarrow p}^{D_2}} = \frac{\frac{1}{2} 9 \cdot 16 \sqrt{\frac{2 \cdot 32}{34}}}{\frac{1}{2} 0 \cdot 57 \sqrt{\frac{32 \cdot 4}{36}}} = \frac{\mu_F^2}{\mu_D^2} \frac{J_H}{J_D} \frac{\alpha}{\alpha'} = \frac{\mu_F^2}{\mu_D^2} \frac{1}{2} \frac{0 \cdot 4}{0 \cdot 7},$$

where  $\alpha$  and  $\alpha'$  denote expressions in the brackets in the theoretical formulæ for  $Z_{p \rightarrow o}^{H_2}$  and  $Z_{o \rightarrow p}^{D_2}$ . We thus obtain for the ratio  $\mu_F^2/\mu_D^2$  a value of about 30 so that the magnetic moment of the diplogen is about 5.5 times smaller than that of the proton being about  $0.2 \cdot 10^{-23}$  C.G.S. units.

In this calculation we have assumed that the value of  $\alpha$  for the collision is the same for both reaction  $p\text{-H}_2 + O_2$  and  $o\text{-D}_2 + O_2$  and that the validity of the theoretical formulæ is retained. It is hoped to confirm this ratio of  $\mu_F/\mu_D$  by comparing the relative efficiencies of other paramagnetic reactions in the two cases, and to estimate the probable difference in  $\alpha$  due to the different zero point vibrations of  $H_2$  and  $D_2$ . In fact one would expect the closest distance of approach  $\alpha$  to be somewhat smaller for  $D_2$  than for  $H_2$ , leading to an even smaller value for the magnetic moment of the diplogen. Stern and Estermann\* came to the same qualitative result in their direct determination of the magnetic moment of the diplogen by molecular ray methods. Thus it is clear that the diplogen has definitely a finite magnetic moment and an abnormal low ratio of  $\frac{\text{magnetic moment}}{\text{mechanical moment}}$ , whereas the proton has an abnormally high one, both at present inexplicable.

Without the unfailing help and encouragement of Professor E. K. Rideal, F.R.S., it would not have been possible for us to carry out this work, and to him we wish to express our most sincere thanks. We are also extremely grateful to Professor Lindemann, F.R.S., for his gift of the liquid hydrogen required. Two of us (A. and L. F.) are glad to express our indebtedness to the Central British Fund for German Jewry and to the Imperial Chemical Industries for financial assistance.

### Summary

It is shown that the ortho-para conversion of diplogen can be catalysed by charcoal and the actual conversion measured by thermal conductivity methods. From the dependence of the ortho-para diplogen equilibrium on temperature

\* 'Z. Physik,' vol. 86, p. 132 (1933).



it is found that for the diplogen nucleus the Bose Einstein statistics are valid, the lowest rotational state being an ortho level, furthermore, that the most probable value of the nuclear spin of diplogen is 1. The heterogeneous ortho-para conversion at 20.4° K proceeds under the experimental conditions at nearly the same rate both for hydrogen and for diplogen. The  $\alpha$ -conversion of ortho-diplogen to normal diplogen by the paramagnetic molecular oxygen is 16 times smaller than the corresponding reaction for hydrogen, indicating that the magnetic moment of the diplogen is about 1/5 of the magnetic moment of the proton.

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## *The Thermal Conductivity of Air*

By T. H. LABY, F.R.S.

(Received September 25, 1933)

When Miss Nelson and the writer prepared in 1929 an article for the 'International Critical Tables' on the thermal conductivity of gases, we found that the value for air had been measured by 19 observers and that the mean departure from the mean was 7%. Further the values obtained by the hot wire method by Weber, Gregory and Archer, and Schneider were higher than the value  $(5.40 \times 10^{-6} \text{ cal cm}^{-1} \text{ sec}^{-1} \text{ deg}^{-1})^*$  which Hertz and the writer had found by a parallel plate method and higher than 13 of the 14 determinations (including hot wire ones) made previous to 1918. In view of these facts it was desirable to repeat the parallel plate method and to obtain evidence as to whether or not there was a systematic difference between the two methods mentioned.

The hot wire method, as used by the experimenters named, has the practical advantages that the temperature of the wire can be kept constant and be accurately measured, and it is convenient and simple. As carried out in the experiments referred to in which fine wires were used it has the disadvantages that the elimination of the convection effects is not attained with certainty, the temperature gradient in the gas is large (which introduces both theoretical and practical difficulties) and there is a temperature discontinuity at the surface of the wire which has to be determined. Hertz and Sutherland have nearly completed in this laboratory a measurement of the thermal conductivity of air with a parallel plate apparatus. This method has the inherent advantages that there are no convection currents in the horizontal lamina of gas used, that the temperature gradient may be made small, and the temperature discontinuity at the surface of the plate is of no significance. In the experiments now in progress radiation is eliminated by using the metal plates at two different separations and considerable improvements have been made as compared with the experiment of Hertz and the writer by the use of a better technique, in the steadiness of the temperatures and in their measurement.

Kannuliuk noticed while the experiment just described was in progress that a development of Kundsen's hot wire method which he had used for measuring the thermal conductivity of the metal of the wire could be used for measuring

\* This and the other values given in this paper for the thermal conductivity of air are for air at 0° C.

gas conductivities. He and Martin in a paper already published and in the one which follows this describe this hot wire method as it is used for measuring gas conductivities. The apparatus is of the simplest construction and the method has an exact theory. As a *thick wire* is used in this apparatus it is free from two of the disadvantages mentioned above, for the temperature gradient can be made small, and the temperature discontinuity is almost or actually negligible. By the use of narrow metal tubes convection effects have been eliminated before molecular conduction begins.

Kannuluk and Martin obtain a value for the conductivity of air of  $5.76 \times 10^{-6}$  cal cm<sup>-1</sup> deg<sup>-1</sup> sec<sup>-1</sup> which is higher than that obtained in the older determinations. Hercus and Sutherland's experiments with the parallel plate method give a value  $5.72 \times 10^{-6}$  which agrees to better than 1% with the value of Kannuluk and Martin.

Thus these two investigations confirm the value\* found by Weber, namely,  $5.74 \times 10^{-6}$ , and show that the parallel plate and certain hot wire methods agree. The accuracy and convenience with which thermal conductivities of gases can be measured appear to have been considerably improved by the method described in the following paper.

We are indebted to Dr. J. K. Roberts for reading the proofs.

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\* Gregory and Archer's value,  $5.83 \times 10^{-6}$ , is not very different.

# *The Thermal Conductivity of some Gases at 0° C*

By W G KANNULUIK, B Sc, and L H MARTIN, Ph D, University of Melbourne

(Communicated by T H Laby, F R S —Received September 25, 1933)

## *Introduction.*

During recent years an increased interest has been displayed in the phenomena of gas conduction, particularly in their application to the observation of molecular changes and chemical dissociations. While relative measurements usually suffice for these purposes, there have also been carefully planned researches on the absolute thermal conductivities of gases, the results of which are of value in the development of the kinetic theory.

A comprehensive account of the methods which have been employed in the past for the measurement of the thermal conductivity of gases is given in a recent paper by Trautz and Zündel,\* who include also a table of the available data to 1931 for air, hydrogen, and carbon dioxide. The lack of agreement between the values obtained by different workers shown in this table can be explained by the smallness of the quantity measured, and by the difficulty of eliminating the heat transfers by convection and by radiation, one or both of which are always present.

The principal methods which have been used in absolute determinations are (1) the parallel plate method, successfully adapted by Hercus and Laby,† and (2) the hot wire method, and its variants, of which the most satisfactory forms are found in the experiments of Weber‡ and of Gregory and Archer,§ and Gregory and Marshall||

The most important work in recent years has been carried out with some form of hot wire method, and the conductivities obtained, notably for air, have been consistently higher than the values obtained with the parallel plate method.

The parallel plate method has many advantages of a fundamental character, and as the hot wire methods have not always been above criticism in matters

\* 'Z. tech. Phys.,' vol 12, p 273 (1931)

† 'Proc. Roy. Soc.,' A, vol 95, p 190 (1919)

‡ 'Ann. Physik,' vol 54, p 325 (1917), vol 82, p 479 (1927)

§ 'Proc. Roy. Soc.,' A, vol 110, p 91 (1926)

|| 'Proc. Roy. Soc.,' A, vol 114, p 354 (1927), vol 118, p 594 (1928)

of detail it was believed that a new series of measurements would be of value if a hot wire method could be developed which met these criticisms satisfactorily. The present paper describes the measurement of the conductivities of a number of gases using a new form of hot wire method in which an exact theory is completely realized with a single simple apparatus.

All hot wire methods are faced with the initial difficulties of eliminating convection, and of making adequate correction for the temperature discontinuity at the wire surface. The problem of separating convection and conduction has been studied by Weber and it is now possible to design a hot wire apparatus in which convection is absent. Briefly, the tube must be vertical, and not too wide. One of the most serious criticisms which can be levelled at recent work is associated with errors introduced by the temperature discontinuity at the wire. In some cases the correction has been neglected, and in others, on account of the small diameter of the wire and the low gas pressures used, the correction has been much larger than is desirable. The temperature discontinuity at the wire surface is related to the accommodation coefficient of the wire for the gas molecules, and it is thus possible to use values of the accommodation coefficients found by other workers in making the necessary correction, but this procedure is not desirable as the coefficient depends on the state of the metal surface. It is preferable to measure the accommodation coefficient with the actual apparatus used for the conductivity experiments, or better, to employ a graphical method, described later, which gives the gas conductivity directly without having to evaluate the accommodation coefficient.

In order to compare the results of different workers, final values of  $k$  are usually reduced to 0° C. In the past this reduction has sometimes involved a considerable extrapolation based on a series of measurements carried out in a relatively short range of temperatures.

In the present experiments the mean temperature of the gas is never greater than 3.3° C and is usually nearer 2° C so that no appreciable error is introduced in reducing the conductivities to 0° C. For this purpose we preferred to employ the temperature coefficients given in the 'International Critical Tables'\* by Laby and Nelson as these are based on experiments made over a wide range of temperature.

The method and tube design are such as to reduce the above sources of error to a minimum. For the majority of gases, there is no convection over a pressure range of some 50 cm. of Hg. The wires used are all so thick that,

\* 'I. C. T.', vol. 5, p. 213

with the possible exception of hydrogen, within this pressure range the correction for the temperature discontinuity can be neglected

### *Method and Theory*

Before describing the method used in these measurements it will be useful to recall the general principles of the hot wire method

A metal wire, which is heated by an electric current, is mounted axially in a glass tube immersed in a constant temperature bath. The wire serves in the dual capacity of a resistance thermometer, and a heater which is at a uniform temperature along a central portion, which will be relatively greater the longer the wire and the smaller its diameter. The lateral heat transfer through the gas is therefore radial only for a limited central portion of the tube. This difficulty has been met in the past by two distinct methods of procedure. In the first, due to Schleiermacher,\* a long thin wire is used and the resistance of a central portion which is at a uniform temperature is measured by introducing two very fine potential leads through the side walls of the tube. In a second method which was devised by Goldschmidt† two tubes are employed, one long and the other short, the differential measurements then refer to a central portion of the longer tube where radial flow conditions hold.

Whichever of these two variants is adopted inconveniently long apparatus are required containing relatively fine wires the mean diameters of which in most experiments have only been determined by weighing methods. As the temperature gradient at the surface of a fine wire is very great it is necessary that it should be exactly circular in cross-section, and of constant diameter, requirements which are best verified by contact methods.

These limitations have been removed in our experiments which are based on the general theory of the distribution of temperature along the wire and in the gas, and makes possible the use of a thick wire mounted in a single short metal tube.

A brief description of the general principles of the present method is as follows. With the ends of the wire and the wall of the tube at  $0^{\circ}\text{C}$  the conductivity of the gas can be calculated from the rate of input of electrical energy, the thermal conductivity of the wire, and its mean temperature. Part of the heat generated is conducted laterally through the gas, while the remainder is dissipated at the ends of the wire. The theory takes into account both of these processes.

\* 'Wied Ann,' vol 34, p. 623 (1888)

† 'Phys Z,' vol 12, p 418 (1911)

With the short thick wires used in this experiment and with gases like air, which are relatively poor conductors, approximately 50% of the total heat generated is dissipated at the ends of the wire. It follows that the accuracy of the gas conductivity determinations depends ultimately on the accuracy with which the thermal conductivity of the wire can be measured. This quantity is measured with the apparatus unchanged in any way, the wire *in situ*, and in the same physical condition as it was when used to measure gas conductivities. The accuracy of this measurement is better than 1 in 300 (see Table II).

*Theory*—A wire of thermal conductivity  $\lambda$  is mounted axially in a tube which is maintained at 0° C. The annular space between wire and tube is filled with a gas of conductivity  $k$ . It follows then that

$$\pi b^2 \lambda \frac{\partial^2 \theta}{\partial z^2} + 2\pi b k \left. \frac{\partial \theta}{\partial r} \right|_{r=b} + I^2 \rho_0 (1 + \alpha \theta) / J = 0, \quad (1)$$

where  $\rho_0 (1 + \alpha \theta)$  is the resistance per unit length of the wire at the temperature  $\theta$ ° C and  $b$  is the radius of the wire.

The first two terms multiplied by  $dz$  represent the net rate of inflow of heat into the element  $dz$  along the wire and over its surface, while the last term multiplied by  $dz$  is the rate of generation of heat in  $dz$ .

If the heat flow from the wire through the gas to the wall of the tube is strictly radial, equation (1) becomes

$$\pi b^2 \lambda \frac{d^2 \theta}{dz^2} + 2\pi b h \theta + I^2 \rho_0 (1 + \alpha \theta) / J = 0, \quad (2)$$

where

$$h = k/b \log_e a/b, \quad (3)$$

and  $a$  is the internal radius of the tube.

The solution\* of equation (2) is as follows

$$\left\{ \left( \frac{1}{\beta l} \right)^2 \left( 1 - \frac{\tanh \beta l}{\beta l} \right) - \frac{2\pi b^2 \lambda J (\bar{R} - R_0)}{R_0^2 I^2 \alpha l} \right\} \quad (4)$$

$$\beta^2 = \frac{2h}{b\lambda} - \frac{I^2 R_0 \alpha}{2\pi b^2 J \lambda}$$

and  $2l$  is the length of the wire inside the tube.

The substitution in this solution of the necessary measured quantities provides an evaluation of  $h$ , which when used with equation (3) yields the gas conductivity  $k$ .

\* Kannuluik, 'Proc Roy Soc,' A, vol 131, p 320 (1931), and Weber, 'Ann. Physik, vol 54, p 165 (1917).

An exact solution of equation (1) takes into consideration the fact that the flow through the gas is not radial, but follows lines of heat flow which show some curvature, particularly at the ends of the tube. The effect of this curvature on the space factor  $b \log_e a/b$  proves to be very small.

The exact solution of equation (1) has already been given in detail in a previous paper,\* and here the final equation only is shown

$$\pi (\bar{R} - R_0)/2R_0\alpha = C_1 N_{01} - 1/3 C_2 N_{03} + 1/5 C_3 N_{05} - \dots, \quad (5)$$

where

$$N_{0n} = I_0(nm)/I_0(np) - K_0(nm)/K_0(np),$$

$n$  being odd, and  $m \equiv \frac{\pi b}{2l}$ ,  $p \equiv \frac{\pi a}{2l}$ , and

$$C_n = \pm 4I^2\rho_0/\pi nJ - \left[ \pi b^2\lambda \left( \frac{n\pi}{2l} \right)^2 - I^2\rho_0\alpha/J \right] N_{0n} - 2\pi b k \left( \frac{n\pi}{2l} \right) N_{1n},$$

positive for  $n = 1, 5, 9, \dots$ ,

negative for  $n = 3, 7, 11, \dots$ ,

with

$$2\pi b N_{1n} = 2\pi b I_1(nm)/I_0(np) N_{0n} + 4l/n I_0(nm) K_0(np),$$

$n$  being odd

The expansions of the Bessel functions  $I_0$ ,  $I_1$ , and  $K_0$  are given in Whittaker and Watson, "Modern Analysis," chap. 17.

The series on the right-hand side of (5) converges rapidly and the calculation of the first three terms allows  $k$  to be obtained by successive approximations.

Equations (4) and (5) have each been used in the calculation of  $k$  for a number of different representative experiments with each tube. In this way the correction is determined, which must be applied to the values of  $k$  deduced from the simpler equation (4). The correction is less than 1%.

Before equation (4) can be used,  $\lambda$  for the wire must be determined. The same apparatus is used to determine both  $\lambda$  and  $k$ . The tube is evacuated to a pressure of the order  $10^{-4}$  mm. Hg, when the lateral loss by molecular conduction is small enough to be neglected.

It has been shown† that  $\lambda$  is given by

$$\lambda = \frac{1}{8} \frac{\bar{R} R_0 I^2 \alpha l}{\pi b^2 J (\bar{R} - R_0)} \left( 1 + \frac{1}{2} \frac{R_0 I^2 \alpha l}{\pi b^2 J \lambda} \right) \left( 1 - \frac{1}{2} \frac{h_p l^2}{b \lambda} \right), \quad (6)$$

\* Kannuluik and Martin, 'Proc. Roy. Soc.', A, vol. 141, p. 144 (1933). We are indebted to Professor T. Cherry for this solution.

† Kannuluik, *loc. cit.*, and Weber, *loc. cit.*



where  $h$ , cal cm<sup>-2</sup> sec<sup>-1</sup> deg<sup>-1</sup> is the small residual lateral loss of heat by radiation. This radiation correction reaches a maximum value in our experiments of 0.5%.

*Description of Apparatus*—We have departed from the usual practice in hot wire work of mounting the wire in a glass tube, preferring with Eucken\* to use metal tubes which have the following advantages —

- (1) The ends of the wire are maintained exactly at the temperature of the bath, the wire being soldered through thin copper caps, which close the tube.
- (2) It is unnecessary to correct for a temperature drop through the wall of the tube, and the gas at the wall is everywhere at the temperature of the bath.
- (3) Eucken† has pointed out that with glass tubes thermal equilibrium is attained very slowly. The use of metal tubes considerably decreases the time taken for the steady state to be reached and it can therefore be detected with greater certainty.

A thorough test of the possible sources of error in the method and apparatus has been made. The space factor has been varied by using two copper tubes of different dimensions, each provided with a copper wire, while the wire material has been varied as well as the space factor, by constructing a third tube of stainless steel provided with a platinum wire. In this way two wires are used, the conductivity of one being more than five times that of the other.

The principles followed in the construction of the tubes may be inferred from the diagram, fig 1, and a detailed description of the third tube.

This tube was of "Two score"‡ stainless steel of internal diameter 1.277 cm and length 10.61 cm. The wall was 1 mm thick. The tube is closed by thin copper caps approximately 1 mm thick, the lower one being insulated from the tube proper by a copper-glass joint. A platinum wire 1.490 mm in



FIG 1

\* 'Phys. Z.', vol 12, p 1002 (1911)

† *Loc. cit.*

‡ Carbon 0.08/0.15, silicon 0.3/0.5, manganese 0.5/0.7, chromium 18/20, nickel 0.15/0.2

diameter is mounted axially in the tube by soldering it through holes in the end caps which were bored accurately central with respect to the tube by rotating this in a lathe. Measurements of the coefficients of linear expansion of different varieties of steel showed that "Two score" had a coefficient of expansion, 0.000010, which is only slightly greater than that of platinum. A platinum wire mounted in a stainless steel tube provides a permanent apparatus, and it was used for measuring the conductivity of a series of different gases.

All three tubes were thus compensated for the expansion which occurs in baths of different temperatures. The dimensions of the copper tubes are shown in the following table -

Table I—Dimensions of Tubes

Apparatus	Length	Internal diameter	Outer diameter	Diameter of wire
	cm	cm	cm	cm
Cu (1)	15.83	0.950	1.526	0.06106
Cu (2)	10.21	1.592	2.207	0.06120
Pt	10.61	1.277	1.472	0.1490

The copper wires for Cu (1) and Cu (2) were drawn from a piece of Hilger's spectroscopically pure copper rod, the platinum wire was a very pure specimen of this metal by Heraeus. The diameters of the wires were measured in terms of standard Johansson gauges using contact methods. Each wire used was circular in section.

The metal tubes were constructed in the following manner. A length of drawn tube, or solid rod through which a hole had been bored, had forced through it a steel ball-bearing the diameter of which was slightly greater than that of the hole.

This produced a smooth central channel which was fairly circular in section but was not quite straight. The tube was pressed as near straight as possible, and was then internally lapped with a series of mild steel laps of increasing diameter, the graded abrasive being held in a spiral groove cut in the lap. The tube was now mounted with a press fit on a hardened steel mandril which had been carefully ground on centres to be straight (as tested against a Brown and Sharp surface plate) and accurately circular in section. The outside was next ground to produce a wall of uniform thickness. Great care was taken in this operation as the outer surface controls the centring of the wire. The tube

was now relapped until it was found that a steel ball fell through it with a slow uniform speed, when the tube was held vertically with one end closed with adhesive tape

A central section of the tube produced in this manner was used in the construction of the apparatus, the diameter being measured by means of a series of "go" and "no go" plug gauges. The internal diameter of the tube was also contoured by dropping into it a ball bearing, the diameter of which was varied, by heating it to different known temperatures. The greatest variation in diameter found in any tube amounted to 0.002 cm.

It may be mentioned here that a slight displacement  $\delta$  of the wire from the axis of the tube does not lead to appreciable error in the space factor. The expression  $b \log_e a/b$  of equation (3) must be replaced by

$$b \log_e \{a/b - a/b \cdot \delta^2/(a^2 - b^2)\},$$

and substitution for particular cases shows that in our experiments the positioning was carried out with more than sufficient accuracy.

The following sources of error have been the most serious in previous hot wire methods\* —

- (1) The transfer of heat through the ends of the wire
- (2) The temperature discontinuity at the surface of the wire
- (3) Heat losses by convection

In our experiments these sources of error are reduced to a minimum. The thermal conductivity  $\lambda$  of the wire is determined with the wire in position. As a result of the temperature discontinuity at the wire and wall surfaces the normal space factor  $b \log_e a/b$  must be replaced by

$$b \{\log_e a/b + \gamma (1/a + 1/b)\},$$

where  $\gamma$  is related to the temperature discontinuity  $\Delta T$  by the equation of Kundt and Warburg

$$\Delta T = -\gamma \frac{dT}{dn}$$

The wires used by us are much thicker than those used by previous workers so that the term containing  $\gamma$  in the space factor is much decreased. Moreover with our apparatus it is unnecessary to work at low gas pressures when  $\gamma$  becomes large, in order to eliminate convection. For example, for air, the gas conductivities are independent of the pressure from 60 to 10 cm Hg. It was considered that the independence of  $k$  of  $p$  over such a wide pressure

\* Weber, 'Ann. Physik,' vol. 82, p. 479 (1927).

range constituted sufficient evidence of the absence of convection in this range. Accommodation coefficients determined at low pressures are in good agreement with existing data, but in obtaining mean values of  $k$ , we have preferred to consider only those values which lie within the pressure range where the effect of the temperature discontinuity has not yet become appreciable.

### *Experimental Procedure*

Apart from measuring the geometrical constants of the wire and tube, the experimental part of the work involves the determination of —

- (1) The electrical constants,  $R_0$  ohm the resistance of the wire at  $0^\circ \text{C}$ , and  $\alpha$  the temperature coefficient of the wire at  $0^\circ \text{C}$
- (2) The thermal conductivity  $\lambda$  of the wire
- (3) The small change in resistance ( $\bar{R} - R_0$ ) ohm of the wire when heated by a suitable current  $I$  amp

As the success of the experiments depends on the accurate determination of a small change in resistance ( $\bar{R} - R_0$ ) corresponding in the majority of cases to a mean rise of temperature of the wire of  $4^\circ$  to  $6^\circ$  only, it is necessary to make very precise determinations of  $R_0$ . For this purpose the pressure of air in the tube is reduced to less than  $10^{-4}$  mm Hg so that the conditions for the application of equation (6) are satisfied. For our tubes the term  $\left(1 + \frac{1}{3} \cdot \frac{R_0 I^2 \alpha l}{\pi b^2 J \lambda}\right)$  in (6) is only slightly greater than unity and may be calculated with ample accuracy using approximate values of  $R_0$  and  $\lambda$ . It is evident from (6) that  $1/\bar{R}$  is a linear function of  $I^2$  multiplied by the factor mentioned above, so that  $1/R_0$  can be obtained by extrapolating  $1/\bar{R}$  to zero current. This method yields very accurate values of  $R_0$ .

To determine  $\alpha$  the tube is filled with hydrogen, and a very small heating current  $I$  is passed through the wire, the resistance of which is determined at the ice point, the steam point and the transition point of sodium sulphate  $32.383^\circ \text{C}$ . Assuming that the resistance of the wire is a quadratic function of the temperature,  $\alpha$  can be determined from the relation

$$R_t = R_0 (1 + \alpha t + \beta t^2)$$

For the copper wires  $\beta$  is positive, for the platinum wire it is negative. While the transition point of sodium sulphate is an excellent fixed point, only material of the highest purity may be used.\* Merck's C.R. was used.

\* See Dickinson and Mueller, 'Bureau of Standards,' vol. 3, p. 655 (1907)

To determine  $\lambda$  the apparatus is evacuated with charcoal and liquid air, and corresponding values of  $\bar{R}$  are obtained for a number of different values of  $I$ , when  $\lambda$  can be calculated from (6). The calculation of the radiation correction has been described in a previous paper\*. In these experiments it amounts to approximately 0.5%, being smaller for the tubes (1) and (2) than for (3).

A three dial type of the "thermo-kraft-frei" potentiometer by Wolff is used to measure  $\bar{R}$  and  $R_0$ , and also the current  $I$  amps in terms of a standard 0.01 ohm resistance and a Weston Standard cell.

After evaluating  $\lambda$ , the additional measurements required to determine  $k$  are a series of corresponding values of  $\bar{R}$  and  $I$  when the tube is filled with a gas at a series of different pressures. The pressures are measured with a U-tube mercury manometer. The apparatus is immersed in a Dewar flask containing pure ice and water which is hand stirred vertically. The approximate values of  $k$  are obtained from equation (4) and these are later reduced so that they agree with the values which would be obtained directly from (5).

The electrical and thermal constants of each of the three wires are given in Table II.

Table II—Constants of Wires

Constants	Cu 1	Cu 2	Pt
$R_0$ (ohm)	0.00864717	0.00547941	0.00601815
$\alpha$	0.004257	0.004244	0.003967
$\beta$	0.042	0.0437	-0.0463
$\lambda$ cal cm <sup>-1</sup> sec <sup>-1</sup> deg <sup>-1</sup>	0.949 ± 0.001	0.929 ± 0.001	0.1872 ± 0.0003

The thermal conductivity  $\lambda$  for Cu 2 is somewhat lower than for Cu 1 although each wire was drawn from the same rod of Hilger's spectroscopically pure copper. This difference is due to different heat treatment, specimens (2) being rapidly cooled after being heated in an annealing furnace. No attempt was made to remove this difference since it provided two copper wires of slightly different conductivity and so gave a check on the method. The mean values of  $\lambda$  are obtained from at least six different corresponding values of  $\bar{R}$  and  $I$ .

Representative runs for air, neon, and hydrogen are shown in Tables III, IV, and V respectively. In these tables, column five, under the heading of  $f$ , gives values of  $(1/\beta I)^2 (1 - \tanh \beta I / \beta I)$ . The values of  $k'$  given in the last

\* Kannuluik, 'Proc Roy Soc.,' A, vol 131, p 320 (1931)

column need to be corrected to the true space factor, and for the temperature of the gas

It will be noticed that the effect of the temperature discontinuity for hydrogen appears at relatively high pressures. For the present purpose it is convenient to divide the phenomenon of gas conduction as a function of the

Table III—Conductivity of dry air measured with tube 1 Mean temperature of gas 3.3° C

Pressure	I	$\bar{R} - R_0$	f	$\beta^*$	$L' \times 10^8$
cm	amps	ohm			
75.3	3 7586 <sub>2</sub>	0 00024024	0 1560	0 04814	5 875
60.1	3 7547 <sub>2</sub>	24052	0 1583	0 04800	5 857
44.0	3 7538 <sub>2</sub>	24041	0 1564	0 04793	5 849
28.6	3 7536 <sub>2</sub>	24040	0 1563	0 04800	5 857
17.3	3 7535 <sub>2</sub>	24052	0 1564	0 04792	5 849
10.3	3 7530 <sub>2</sub>	24036	0 1564	0 04792	5 849
5.0	3 7516 <sub>2</sub>	24058	0 1566	0 04782	5 836
0.9	3 7504 <sub>2</sub>	24232	0 1579	0 04712	5 750

Mean  $L'$  in pressure range 60 to 10 cm  $5.85_2 \pm 0.007 \times 10^{-8}$  at 3.3° C Mean  $k'$  at 0° C, using a temperature coefficient  $0.003, 5.793 \times 10^{-9}$ . On using correct space factor,  $L_0 = 5.748 \times 10^{-8}$  cal cm<sup>-1</sup> sec<sup>-1</sup> deg<sup>-1</sup>

Table IV—Conductivity of pure neon measured with tube 3 Mean temperature of gas 1.7° C

Pressure	I	$\bar{R} - R_0$	f	$\beta^*$	$L' \times 10^8$
cm	amps	ohm			
67.0	4 3289 <sub>2</sub>	0 00008918	0 1503	0 1132	11.28
67.0	4 3260 <sub>2</sub>	8795	0 1503	0 1132	11.28
55.6	4 3261 <sub>2</sub>	8788	0 1502	0 1133	11.29
42.4	4 3263 <sub>2</sub>	8791	0 1502	0 1133	11.29
30.6	4 3264 <sub>2</sub>	8785	0 1501	0 1134	11.30
19.4	4 3263 <sub>2</sub>	8785	0 1501	0 1134	11.30
19.4	3 5473 <sub>2</sub>	5870	0 1492	0 1135	11.31*

\* Mean temperature 1.1° C

Mean  $k'$  in pressure range 67.0 to 19.4 cm,  $11.29 \pm 0.01 \times 10^{-8}$  at 1.7° C Mean  $k'$  at 0° C using a temperature coefficient  $0.003, 11.23 \times 10^{-9}$ . Using correct space factor  $L_0 = 11.12 \times 10^{-8}$  cal cm<sup>-1</sup> sec<sup>-1</sup> deg<sup>-1</sup>

pressure into two regimes. At relatively high pressures the conductivity is independent of pressure, while at moderately low pressures there is a range determined by the geometry of the apparatus and the mean free path of the gas where the effect of the temperature discontinuity must be taken into account.

This is accomplished by replacing the normal temperature gradient  $\theta/d$  by  $\theta/(d + \Delta)$  where  $\Delta$  is inversely proportional to the pressure. If this correction be introduced into the space factor of the apparatus it can be shown\* that  $p/k_0$  is proportional to  $p$  down to pressures at which molecular conduction appears. If  $p/k_0$  and  $p$  are plotted using all the data in Table V, a straight line is obtained from which the conductivity is deduced from the cotangent of the angle made with the pressure axis. The value so obtained,  $k' = 41.85 \times 10^5$ , is in excellent agreement with the value given above. This procedure does not involve a knowledge of the accommodation coefficient itself.

Table V—Conductivity of pure hydrogen measured with tube 2. Mean temperature of gas 0.9° C

Pressure	I	$\bar{R} - R_s$	$f$	$\beta^2$	$k \times 10^5$
cm	amps	ohm			
63.1	4.2315 <sub>8</sub>	0.00004280	0.08414	0.2957	41.82
52.6	4.2217 <sub>8</sub>	4280	8432	0.2951	41.74
38.8	4.2221 <sub>8</sub>	4284	8402	0.2963	41.90
31.2	4.2318 <sub>8</sub>	4298	8408	0.2961	41.87
21.8	4.2349 <sub>8</sub>	4325	8453	0.2940	41.57
10.5	4.2368 <sub>8</sub>	4362	8515	0.2911	41.16
5.3	4.2371 <sub>8</sub>	4402	8592	0.2875	40.65
2.6	4.2374 <sub>8</sub>	4729	9230	0.2617	37.01

Mean  $k'$  in pressure range 63 to 31 cm,  $41.83 \pm 0.05 \times 10^{-5}$  at 0.9° C. Mean  $k'$  at 0° C, using a temperature coefficient  $0.003 = 41.71 \times 10^{-5}$ . On using correct space factor,  $k_s = 41.3 \times 10^{-5}$  cal sec<sup>-1</sup> cm<sup>-1</sup> deg<sup>-1</sup>.

It has already been mentioned that a difficulty arises in reducing  $k$  to 0° C on account of some uncertainty in the values of the temperature coefficients of  $k$ . The correction is a small one in this work and it has been sufficient to use one significant figure in the values of the coefficients adopted.

### Experimental Results

The methods employed in the preparation of pure gases are those recommended in Traver's "Study of Gases" and in the "Handbuch der Arbeits Methoden in der Anorganischen Chemie," vol. 4.

**Air**—The air was dried by passing it through phosphorus pentoxide. The carbon dioxide was not removed, as Weber has shown that its presence does not appreciably affect the conductivity of air. The results for air are shown in Table VI. The values of  $k'$  in column 3 are the mean values obtained with each tube, the corresponding gas temperatures being shown in column 2. Column 5 gives the conductivity corrected for temperature, and column 6 the final value for each tube after a correction has been made for the curvature of

\* Gregory and Aroher, 'Phil. Mag.', vol. 15, p. 301 (1933), see also Kannuluik and Martin, 'Proc. Roy. Soc. A', vol. 141, p. 144 (1933).

the lines of heat flow This description also applies to further tables giving results for other gases

Table VI—Air

Apparatus	Mean temperature of gas	$l' \times 10^3$	Temperature coefficient	$k' \times 10^3$ at 0° C	$k_s \times 10^{13}$
	° C				
Cu 1	3.3	$5.85_s \pm 0.007$	0.003	5.79	5.75
Cu 2	1.7	$5.83_s \pm 0.009$	0.003	5.81	5.76
Pt	1.7	$5.84_s \pm 0.008$	0.003	5.82	5.76

Mean value of  $k_s = 5.76 \times 10^{-8}$  cal cm<sup>-1</sup> sec<sup>-1</sup> deg<sup>-1</sup>

The agreement between the three tubes is satisfactory and is certainly much better than we had hoped to obtain when planning these experiments. We believe that the absolute accuracy is somewhat less than is indicated by this table, and is probably better than 0.5%.

**Hydrogen**—The apparatus was first pumped to an X-ray vacuum, and then filled with pure hydrogen which was prepared by passing commercial hydrogen through a palladium tube heated electrically to a bright red. The whole apparatus was washed out several times with hydrogen so prepared.

Table VII—Hydrogen

Apparatus	Mean temperature of gas	$l' \times 10^3$	Temperature coefficient	$l' \times 10^3$ at 0° C	$k_s \times 10^3$
Cu 1	1.3	$41.83 \pm 0.10$	0.003	41.66	41.3
Cu 2	0.9	$41.83 \pm 0.05$	0.003	41.71	41.3
Pt	1.7	$41.71 \pm 0.07$	0.003	41.52	41.2

Mean value of  $k_s = 41.3 \times 10^{-8}$  cal cm<sup>-1</sup> sec<sup>-1</sup> deg<sup>-1</sup>

**Carbon Dioxide**—Two methods were used to produce pure carbon dioxide. For tube 1, the gas was produced by heating pure magnesium carbonate, the gas being passed through concentrated sulphuric acid, and then through phosphorus pentoxide. For tube 2, pure gas was obtained by Lord Rayleigh's method, in which the gas is drawn off an inverted cylinder after a considerable volume has been blown off from the cylinder in the erect position. The gas was dried by passing it through phosphorus pentoxide. As for all gases, the apparatus was first pumped to an X-ray vacuum, and was then washed out several times with the gas, before being filled.



Table VIII—Carbon Dioxide

Tube	Mean temperature of gas	$k' \times 10^4$	Temperature coefficient	$k' \times 10^4$ at 0° C	$k_s \times 10^4$
Cu 1	2.4	$3.492 \pm 0.024$	0.004	3.46	3.44
Pt	2.0	$3.508 \pm 0.008$	0.004	3.48	3.43

Mean value of  $k_s = 3.43 \times 10^{-4}$  cal cm<sup>-1</sup> sec<sup>-1</sup> deg<sup>-1</sup>

*Oxygen*—The gas was prepared by heating potassium permanganate and was then passed through concentrated sulphuric acid and over phosphorus pentoxide

Table IX—Oxygen

Tube	Mean temperature of gas	$k \times 10^4$	Temperature coefficient	$k \times 10^4$ at 0° C	$k_s \times 10^4$
	° C				
Cu 1	2.7	$5.929 \pm 0.006$	0.003	5.88	5.84
Pt	1.7	$5.913 \pm 0.009$	0.003	5.88	5.82

Mean value of  $k_s = 5.83 \times 10^{-4}$  cal cm<sup>-1</sup> sec<sup>-1</sup> deg<sup>-1</sup>

In view of the excellent agreement in the results obtained with the different tubes for air, hydrogen, carbon dioxide, and oxygen, the conductivities of the remaining gases, carbon monoxide, nitrous oxide, and the inert gases, helium, neon, and argon have been measured with the platinum wire tube only

*Carbon Monoxide and Nitrous Oxide*—Carbon monoxide was obtained by the interaction of pure concentrated sulphuric acid and pure sodium formate. The carbon dioxide present was removed by passing the gas through caustic potash, and the gas was finally dried by passing it through phosphorus pentoxide

Nitrous oxide was generated by heating pure ammonium nitrate to a temperature not greater than 250° C, thus preventing the formation of nitric oxide. The gas was passed over caustic potash and then through concentrated sulphuric acid and phosphorus pentoxide

*Monatomic Gases*—The helium and neon used in this work were obtained from the British Oxygen Company, and the argon from Philips Lamp Company (Holland). The certificates supplied with these gases gave the neon and argon as 100% pure and the helium 100% pure except for a trace of neon. The conductivities of helium were determined for the gas as received and after it

Table X—Carbon Monoxide and Nitrous Oxide

Gas	Mean temperature of gas	$k' \times 10^8$	Temperature coefficient	$k' \times 10^8$ at 0° C	$k_s \times 10^8$
Carbon monoxide	° C 1 7	5 447 ± 0 008	0 003	5 420	5 37
Nitrous oxide	2 0	3 677 ± 0 006	0 004	3 645	3 61

had been passed through charcoal immersed in liquid air. It was found that the second value was approximately 2% greater than the first, and this is probably due to the adsorption by the charcoal of the trace of neon stated to be present. The same procedure was adopted with neon, but here no change was observed in the conductivity.

It was also our intention to pass the argon over heated calcium to eliminate possible impurities, but on account of an accident to the apparatus this has been postponed to a further series of experiments on temperature coefficients. However, the conductivity of the argon as received has been measured, and comparison of the value obtained with the values of other observers seems to imply the absence of any appreciable impurity, as this would certainly lead to a higher value.

Table XI—Inert Gases

Gas	Mean Temperature of gas	$k' \times 10^8$	Temperature coefficient	$k' \times 10^8$ at 0° C	$k_s \times 10^8$
Helium	2 0	34 83 ± 0 02	0 003	34 62	34 30
Neon	1 7	11 29 ± 0 01	0 003	11 23	11 12
Argon	2 0	3 88 ± 0 009	0 003	3 86	3 82

### Discussion of Results

Table XII contains the thermal conductivities at 0° C obtained by the authors, together with those for the same gases obtained by Weber and by Gregory, Archer, and Marshall, all of whom have carried out extensive series of very careful measurements using some form of the hot wire method. Our values appear to be in fair agreement with those given by Weber, particularly with his more recent values for air, carbon dioxide, and neon. There is some divergence, however, between our values and those of Gregory, Archer, and Marshall, who obtain values which are, for some gases, 2% or 3% greater than ours. Unfortunately serious errors may easily be introduced in reducing

Table XII— $k_0 \times 10^5 \text{ cal sec}^{-1} \text{ deg}^{-1} \text{ cm}^{-1}$ 

Gas	K and M	Weber		G A and M
		1917	1927	
Air	5.76	5.68	5.740	5.85
H <sub>2</sub>	41.3	41.65	—	40.43
CO <sub>2</sub>	3.43	3.393	3.431	3.60
O <sub>2</sub>	5.83	5.768	—	5.89
CO	5.37	—	—	5.63
N <sub>2</sub> O	3.61	3.530	—	3.74
He	34.3	34.38	—	—
Ne	11.12	10.89	11.04	—
Ar	1.82	1.850	—	—

conductivities to 0° C. For example, with air, Gregory and Archer studied the conductivity over a temperature range 7.4° C to 12.1° C and in reducing the conductivity to 0° C were obliged to extrapolate some 7°. The value of the temperature coefficient implied in this extrapolation was 0.00297. Weber has studied the variation of the conductivity of air over a range of temperatures extending from 3° C to 18.5° C and finds a temperature coefficient of 0.00365, which he points out if used to reduce the data of Gregory and Archer leads to a value for the conductivity at 0° C of  $5.76 \times 10^{-5}$ . This value agrees with that found by Weber and ourselves. It may be added, however, that the value of the temperature coefficient used by Gregory and Archer is in excellent agreement with the value given by Laby and Nelson in the International Critical Tables.

In the single experiment on hydrogen, Gregory and Archer give a smaller conductivity than either Weber or ourselves. An examination of the results of Gregory and Archer lead us to suppose that they have not made a sufficient correction for the temperature "discontinuity," and that when the necessary correction is applied, their result is in excellent agreement with ours.

The value found by Weber for this gas is nearly 1% greater than ours. As Weber in his purification process passed the gas slowly over charcoal immersed in liquid air, this discrepancy may be accounted for by the presence in his hydrogen of a greater proportion of para-hydrogen than is typical of the ordinary gas.

It has been shown\* that ordinary hydrogen, which is a mixture of ortho- and para-hydrogen present in the ratio of 3 to 1, if allowed to remain in contact

\* Bonhoeffer and Harteck, 'Z. phys. Chem.', B, vol. 4, p. 113 (1929), Eucken and Hiller, 'Z. phys. Chem.', B, vol. 4, p. 158 (1929), Dennison, 'Proc. Roy. Soc.', A, vol. 115, p. 483 (1927).

with charcoal at the temperature of liquid air suffers a transformation, the equilibrium mixture now being one in which the ratio of ortho- to para- is 1 2 1. This mixture is quite stable and persists for a considerable time after the gas has been raised again to room temperature. The specific heat of the ordinary 3 to 1 mixture is 2.44R, while that of the mixture characteristic of the liquid air temperature is 2.49R, and since the thermal conductivity is proportional to the specific heat, a 2% increase in conductivity might be expected as a result of the change at the low temperature.

While it is probable that in Weber's purification process the gas was moving too quickly through the charcoal to be transformed to the equilibrium mixture, Senftleben† has shown that a very marked change (sufficiently great to be easily demonstrated to a large audience) takes place even in a minute or two, and it might therefore be reasonably expected that Weber's value will be higher than the real value for ordinary hydrogen.

*Maxwell's Relation*  $k_0 = \eta_0 c_v$ .—Table XIII gives values of  $k_0$ , the viscosity  $\eta$ , and  $c_v$ , while the fifth column shows the constant  $\epsilon$  for the different gases calculated from

$$\epsilon = k_0 / \eta_0 c_v$$

The values obtained for  $\epsilon$  for helium, argon, and neon agree well with the value 2.5 deduced theoretically for monatomic gases by Maxwell, Chapman, and Enskog‡

Table XIII

Gas	$k_0 \times 10^8$	$\eta_0 \times 10^8$	$c_v$	$\epsilon$	$\frac{1}{2}(\gamma - 1)$
He	34.3	18.76	0.746	2.45	2.44
Ne	11.12	29.81	(0.150)*	(2.50)	2.44
Ar	3.82	21.02	0.0745	2.44	2.44
H <sub>2</sub>	41.3	8.60	2.43	2.00	1.90
Air	5.76	17.22	0.171	1.96	1.91
O <sub>2</sub>	5.83	19.31	0.167	1.92	1.95
CO	5.37	16.65	0.178	1.81	1.91
CO <sub>2</sub>	4.43	13.74	0.153	1.64	1.72
N <sub>2</sub> O	3.61	13.66	0.155	1.71	1.73

$c_v$  values: Partington and Shilling, 'Specific Heats of Gases,' p. 201 (1924), Landolt and Bornstein, 'Supp.,' vol. 2, Table 264A, p. 1227 (1931)

$\eta_0$  values: Landolt and Bornstein (loc. cit.)

\*  $c_v$  calculated from  $\frac{1}{2}R$

† 'Z. phys. Chem.,' B, vol. 4, p. 169 (1929)

‡ Jeans, "The Dynamical Theory of Gases," chap. 12

Eucken has suggested a formal combination of Chapman's expression  $k_0 = 5/2 \eta_0 c_v$  and the well-known equation  $c_v = 3/2 (1 + \beta) R/Jm$  for polyatomic gases. The sixth column contains the values of  $\frac{1}{2} (9\gamma - 5)$  which according to this method of introducing the rotational energy of polyatomic molecules should equal  $\epsilon$ .

In conclusion, we desire to express our appreciation of the continued interest Professor Laby has taken in this work. We are indebted to him for many helpful suggestions and for placing every facility at our disposal. We wish to thank Mr. A. H. Turner, of the Commonwealth Radium Laboratory, for help in the manipulation of the inert gases. One of us (W. G. K.) is indebted to the Melbourne University for a Fred Knight scholarship which enabled him also to take part in this work.

We are also indebted to Dr. J. K. Roberts for reading the proofs.

### Summary

The thermal conductivities of the gases, air,  $H_2$ ,  $CO_2$ ,  $O_2$ , CO,  $N_2O$ , He, Ne, and Ar, have been measured using a hot wire method. The method is based on an exact theory for the flow of heat from an electrically heated short thick wire mounted axially in a metal tube maintained at a uniform temperature.

The apparatus realizes completely the requirements of the theory, is simple in construction and rapid in working. Measurements carried out on several gases with three separate pieces of apparatus show excellent agreement. These differ not only in their dimensions, but also in the material of the axial wire.

Possible sources of error in hot wire methods generally are discussed, and our results are compared with previous determinations for the above gases by other workers.

The constant  $\epsilon$  in the kinetic theory relation  $k_0 = \epsilon \eta_0 c_v$  is evaluated for the above gases. The values of  $\epsilon$  obtained for He, Ne, and Ar are in good agreement with Chapman's theoretical value 2.5 for a monatomic gas.

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## *The Calculation of Wave Resistance*

By T H HAVELOCK, F R S

(Received January 25, 1934 )

1 The wave resistance of a body moving in a frictionless liquid has been calculated by various methods. In a few cases it has been found directly as the resultant of the fluid pressures on the surface of the body. Another method, which has been more generally useful, involves the introduction of a certain type of fluid friction into the equations of motion. The wave resistance is then found by calculating the rate of dissipation of energy and taking the limiting value when the frictional coefficient is made vanishingly small. This method has certain important analytical advantages, nevertheless it is highly artificial. A third method, dealing directly with a frictionless liquid, consists in examining the flow of energy in the wave motion, this has hitherto been used only for two-dimensional problems when the wave motion consists of simple waves with straight parallel crests, the usual theory of group velocity being directly applicable.

In the following note this method is extended to three-dimensional fluid motion. Although no new special results are obtained so far as expressions for wave resistance are concerned, it seemed of sufficient interest to obtain them by this direct method, namely, by considering the flow of energy and the rate of work across planes far in advance and far in the rear of the moving body.

These quantities are examined first for a free wave pattern of simple type. Then a general expression is given for wave resistance in terms of the velocity potential of the free wave pattern to which the disturbance approximates at a great distance in the rear, and this is applied to a general form of wave pattern and to some special cases. Finally, a similar examination is made of a certain problem when the water is of finite depth.

2 With the origin  $O$  in the free surface of deep water, and  $Oz$  vertically upwards, the surface condition is

$$\frac{\partial^2 \phi}{\partial t^2} - g \frac{\partial \zeta}{\partial t} = 0, \quad z = 0, \quad (1)$$

where  $\phi$  is the velocity potential and  $\zeta$  the surface elevation. For a wave

pattern advancing steadily with velocity  $c$  in the direction  $Ox$ , we may write (1) in the form

$$\frac{\partial^2 \phi}{\partial x^2} + \kappa_0 \frac{\partial \phi}{\partial x} = 0, \quad z = 0, \quad (2)$$

with  $\kappa_0 = g/c^2$

A simple plane wave advancing in a direction making an angle  $\theta$  with  $Oz$  is given by

$$\left. \begin{aligned} \zeta &= a \sin \{ \kappa_0 \sec^2 \theta (x \cos \theta + y \sin \theta - ct \cos \theta) \} \\ \phi &= ac \cos \theta e^{i\kappa_0 \sec^2 \theta} \cos \{ \kappa_0 \sec^2 \theta (x \cos \theta + y \sin \theta - ct \cos \theta) \} \end{aligned} \right\} \quad (3)$$

We may generalize this to obtain a free wave pattern made up of plane waves advancing in all directions, so that the pattern itself moves steadily with velocity  $c$  in the direction  $Ox$ , we have then

$$\zeta = \int_{-\pi}^{\pi} f(\theta) \sin \{ \kappa_0 \sec^2 \theta (x \cos \theta + y \sin \theta - ct \cos \theta) \} d\theta \quad (4)$$

We shall suppose in the first place that the pattern is symmetrical with respect to  $Ox$ , so that we have

$$\left. \begin{aligned} \zeta &= 2 \int_0^{\pi} f(\theta) \sin (\kappa_0 x' \sec \theta) \cos (\kappa_0 y \sin \theta \sec^2 \theta) d\theta \\ \phi &= 2c \int_0^{\pi} f(\theta) e^{i\kappa_0 x' \sec \theta} \cos (\kappa_0 x' \sec \theta) \cos (\kappa_0 y \sin \theta \sec^2 \theta) \cos \theta d\theta \end{aligned} \right\}, \quad (5)$$

with  $x' = x - ct$

Consider a fixed vertical plane  $x = \text{constant}$ . The rate of flow of total energy across this plane is given by

$$\frac{1}{2} \rho c \int_{-\infty}^{\infty} dz \int_{-\infty}^{\infty} \left\{ \left( \frac{\partial \phi}{\partial x} \right)^2 + \left( \frac{\partial \phi}{\partial y} \right)^2 + \left( \frac{\partial \phi}{\partial z} \right)^2 \right\} dy + \frac{1}{2} g \rho c \int_{-\infty}^{\infty} \zeta^2 dy \quad (6)$$

The variable part of the fluid pressure being  $\rho \partial \phi / \partial t$ , or  $-\rho c \partial \phi / \partial x$ , the rate at which work is being done across the same plane is

$$\rho c \int_{-\infty}^{\infty} dz \int_{-\infty}^{\infty} \left( \frac{\partial \phi}{\partial x} \right)^2 dy \quad (7)$$

We shall assume that the wave pattern is such that these quantities are finite and determinate

To evaluate these expressions with the values (5) for  $\phi$  and  $\zeta$  we use the following theorem

If

$$\left. \begin{aligned} F_1(y) &= \int_0^\infty (A_1 \cos yu + B_1 \sin yu) du, \\ F_2(y) &= \int_0^\infty (A_2 \cos yu + B_2 \sin yu) du, \end{aligned} \right\} \quad (8)$$

A, B, being functions of  $u$ , then

$$\int_{-\infty}^{\infty} F_1(y) F_2(y) dy = \pi \int_0^\infty (A_1 A_2 + B_1 B_2) du, \quad (9)$$

assuming that the integrals are convergent

To take one of the integrals in (6) as an example we have

$$\frac{\partial \phi}{\partial x} = -2\kappa_0 c \int_0^{1\pi} f(\theta) e^{\kappa_0 x' \sec \theta} \sin(\kappa_0 x' \sec \theta) \cos(\kappa_0 y \sin \theta \sec^2 \theta) d\theta \quad (10)$$

To put this into the form (8), we write  $u = \kappa_0 \sin \theta \sec^2 \theta$ , then carry out the process (9) and finally replace the variable  $u$  in terms of  $\theta$ , it is clear that we shall have to introduce into the integral in the final form a factor  $d\theta/du$ , that is, a factor  $\cos^3 \theta / \kappa_0 (1 + \sin^2 \theta)$ . Thus we have

$$\begin{aligned} \int_{-\infty}^{\infty} dz \int_{-\infty}^{\infty} \left( \frac{\partial \phi}{\partial x} \right)^2 dy \\ &= 4\pi \kappa_0 c^2 \int_{-\pi}^{\pi} dz \int_0^{1\pi} \{f(\theta)\}^2 e^{2\kappa_0 x' \sec \theta} \sin^2(\kappa_0 x' \sec \theta) \frac{\cos^3 \theta d\theta}{1 + \sin^2 \theta} \\ &= 2\pi c^2 \int_0^{1\pi} \{f(\theta)\}^2 \sin^2(\kappa_0 x' \sec \theta) \frac{\cos^3 \theta d\theta}{1 + \sin^2 \theta} \end{aligned} \quad (11)$$

From (6) we find in this way that the rate of flow of total energy across the vertical plane is

$$\begin{aligned} \pi \rho c^2 \int_0^{1\pi} \{f(\theta)\}^2 \{ (3 - \sin^2 \theta) \sin^2(\kappa_0 x' \sec \theta) \\ + (1 + \sin^2 \theta) \cos^2(\kappa_0 x' \sec \theta) \} \frac{\cos^3 \theta d\theta}{1 + \sin^2 \theta}, \end{aligned} \quad (12)$$

and that the rate at which work is being done across this plane is

$$2\pi \rho c^2 \int_0^{1\pi} \{f(\theta)\}^2 \sin^2(\kappa_0 x' \sec \theta) \frac{\cos^3 \theta d\theta}{1 + \sin^2 \theta} \quad (13)$$

It is the difference of these two quantities that is significant for our purpose, it is, as would be expected, independent of the time and of the position of the



plane Subtracting (13) from (12), we find that the rate at which energy is being propagated less the rate of work reduces to the simple expression

$$\pi \rho c^3 \int_0^{1\pi} \{f(\theta)\}^2 \cos^3 \theta \, d\theta \quad (14)$$

It may be noted that if we take mean values of (12) and (13) we have as the mean rate of flow of energy

$$2\pi \rho c^3 \int_0^{1\pi} \{f(\theta)\}^2 \frac{\cos^3 \theta}{1 + \sin^2 \theta} \, d\theta, \quad (15)$$

and as the mean rate of work

$$\pi \rho c^3 \int_0^{1\pi} \{f(\theta)\}^2 \frac{\cos^3 \theta}{1 + \sin^2 \theta} \, d\theta \quad (16)$$

The connection indicated in (15) and (16) is a generalization of the well-known result for simple plane waves that the mean rate of work is half the mean rate of flow of energy

3 Consider now the forced wave pattern produced by a body moving through the liquid, or by a localized pressure disturbance The complete surface elevation may be separated into a local disturbance and a wave pattern In a frictionless liquid a possible solution is one in which the wave pattern extends to an infinite distance in advance of the body as well as in the rear The determinate practical solution is that for which the wave pattern vanishes at a great distance in advance, and we may suppose this obtained by superposing over the whole surface a suitable free wave pattern In that case, considering the flow of energy and rate of work across two fixed vertical planes, one far in advance and the other far in the rear, we see that (14) is equal to  $Rc$ , where  $R$  is the wave resistance Hence we have

$$R = \pi \rho c^3 \int_0^{1\pi} \{f(\theta)\}^2 \cos^3 \theta \, d\theta, \quad (17)$$

when the wave pattern at a great distance to the rear approximates to the form (4)

For example, the forced wave pattern produced by a submerged sphere, or more precisely by a horizontal doublet of moment  $M$  at depth  $f$ , approximates at a great distance behind the disturbance, to the free wave pattern

$$\zeta = \frac{4\kappa_0^3 M}{c} \int_{-\pi}^{1\pi} \sec^4 \theta \, e^{-\kappa_0 f \sec \theta} \sin \{\kappa_0 (x' \cos \theta + y \sin \theta) \sec^2 \theta\} \, d\theta \quad (18)$$

Hence, from (17), the wave resistance is

$$R = 16\pi\rho\kappa_0^4 M^2 \int_0^{1\pi} \sec^3 \theta e^{-2\kappa_0 f \sec^3 \theta} d\theta, \quad (19)$$

which is the known result for this case

4 Before generalizing these results we may put (6) and (7) into an explicit form for the wave resistance

The kinetic energy of the liquid in a strip between two parallel vertical planes at a distance  $\delta x$  apart is

$$\frac{1}{2}\rho \delta x \int_{-\infty}^{\infty} dz \int_{-\infty}^{\infty} \left\{ \left( \frac{\partial \phi}{\partial x} \right)^2 + \left( \frac{\partial \phi}{\partial y} \right)^2 + \left( \frac{\partial \phi}{\partial z} \right)^2 \right\} dy \quad (23)$$

Transform (23) into the equivalent form of a surface integral over the boundaries of this portion of fluid, assuming the wave pattern to be such that the various integrals are convergent. Thus we obtain the rate of flow of kinetic energy across a vertical plane as

$$\frac{1}{2}\rho \int_{-\infty}^{\infty} \left( \phi \frac{\partial \phi}{\partial z} \right)_{z=0} dy + \frac{1}{2}\rho c \int_{-\infty}^{\infty} dz \int_{-\infty}^{\infty} \left\{ \phi \frac{\partial^2 \phi}{\partial x^2} + \left( \frac{\partial \phi}{\partial x} \right)^2 \right\} dy \quad (24)$$

Further, we may transform the other terms in (6) and (7) by using the surface condition (2) together with  $g\zeta = -c\partial\phi/\partial x$  at  $z=0$

Finally, equating the difference between (6) and (7) to  $Rc$ , we obtain for the wave resistance

$$R = \frac{\rho}{2\kappa_0} \int_{-\infty}^{\infty} \left\{ \left( \frac{\partial \phi}{\partial x} \right)^2 - \phi \frac{\partial^2 \phi}{\partial x^2} \right\}_{z=0} dy - \frac{1}{2}\rho \int_{-\infty}^{\infty} dz \int_{-\infty}^{\infty} \left\{ \left( \frac{\partial \phi}{\partial x} \right)^2 - \phi \frac{\partial^2 \phi}{\partial x^2} \right\} dy \quad (25)$$

In this expression  $\phi$  is the velocity potential of the free wave pattern to which the disturbance approximates at a great distance in the rear. Considering the disturbance produced by a body of any form, it appears that this free wave pattern must be expressible, in general, in the form

$$\begin{aligned} \zeta = & \int_{-1\pi}^{1\pi} f(\theta) \sin \{ \kappa_0 \sec^2 \theta (x' \cos \theta + y \sin \theta) \} d\theta \\ & + \int_{-1\pi}^{1\pi} F(\theta) \cos \{ \kappa_0 \sec^2 \theta (x' \cos \theta + y \sin \theta) \} d\theta, \end{aligned} \quad (26)$$

that is, in the form

$$\zeta = \int_0^{1\pi} (P_1 \sin A \cos B + P_2 \cos A \sin B + P_3 \cos A \cos B + P_4 \sin A \sin B) d\theta, \quad (27)$$

where  $A = \kappa_0 x' \sec \theta$ ,  $B = \kappa_0 y \sin \theta \sec^2 \theta$

The corresponding velocity potential is

$$\phi = c \int_0^{1\pi} (P_1 \cos A \cos B - P_2 \sin A \sin B - P_3 \sin A \cos B + P_4 \cos A \sin B) e^{i\kappa_0 \sec^2 \theta} \cos \theta \, d\theta \quad (28)$$

With this value of  $\phi$  in (25), we use (8) and (9) to evaluate the integrations with respect to  $y$  as in § 2, and we obtain readily the general result

$$R = \frac{1}{2} \pi \rho c^3 \int_0^{1\pi} (P_1^2 + P_2^2 + P_3^2 + P_4^2) \cos^3 \theta \, d\theta \quad (29)$$

The actual calculation of the quantities  $P$  for a body of given form is, of course, another problem. Methods in use at present amount to replacing the body by some approximately equivalent system of sources and sinks, the functions  $P$  then appear, in general, in the form of integrals taken over the surface of the body. We need not consider these here as the expressions for  $R$  given above lead to the same results as those obtained previously by different methods.

§ It is of interest to examine a similar problem when the water is of finite depth  $h$ . It is clear from the derivation of (25) that we may use it in this case also, taking the lower limit of integration with respect to  $z$  to be  $-h$  instead of  $-\infty$ .

For the simple symmetrical type of free wave pattern given by (4), the corresponding velocity potential is

$$\phi = 2c \int_0^{1\pi} f(\theta) \frac{\cosh \kappa(z+h)}{\sinh \kappa h} \cos(\kappa x' \cos \theta) \cos(\kappa y \sin \theta) \cos \theta \, d\theta, \quad (30)$$

the relation between  $\kappa$  and  $\theta$  being

$$\kappa - \kappa_0 \sec^2 \theta \tanh \kappa h = 0 \quad (31)$$

We shall assume first  $\kappa_0 h > 1$ , that is  $c^2 < gh$ , so that (31) as an equation for  $\kappa$  has one real root for each value of  $\theta$  in the range of integration. In evaluating (25) we carry out the integrations with respect to  $y$  by means of (8) and (9). For this we have to change from an integration in  $\theta$  to one in a variable  $u$  given by

$$u = \kappa \sin \theta, \quad (32)$$

together with (31). The corresponding factor  $d\theta/du$  has now the value

$$\frac{\cos^2 \theta (\coth \kappa h - \kappa h \operatorname{cosech}^2 \kappa h)}{\kappa_0 (1 + \sin^2 \theta - \kappa_0 h \operatorname{sech}^2 \kappa h)} \quad (33)$$

We have, for example,

$$\begin{aligned} & \int_{-h}^0 dz \int_{-\infty}^{\infty} \left( \frac{\partial \phi}{\partial x} \right)^2 dy \\ &= 4\pi c^2 \int_{-h}^0 dz \int_0^{1\pi} \{f(\theta)\}^2 \frac{\cosh^2 \kappa(z+h) (\coth \kappa h - \kappa h \operatorname{cosech}^2 \kappa h)}{\kappa_0 \sinh^2 \kappa h (1 + \sin^2 \theta - \kappa_0 h \operatorname{sech}^2 \kappa h)} \\ & \quad \times \sin^2 (\kappa x' \cos \theta) \kappa^2 \cos^2 \theta d\theta \\ &= 2\pi c^2 \int_0^{1\pi} \{f(\theta)\}^2 \frac{\coth^2 \kappa h - \kappa^2 h^2 \operatorname{cosech}^4 \kappa h}{\kappa_0 (1 + \sin^2 \theta - \kappa_0 h \operatorname{sech}^2 \kappa h)} \sin^2 (\kappa x' \cos \theta) \kappa \cos^2 \theta d\theta \end{aligned} \quad (34)$$

Evaluating the remaining terms in (25), we obtain after a little reduction the result

$$R = \pi \rho c^2 \int_0^{1\pi} \{f(\theta)\}^2 (\coth \kappa h - \kappa h \operatorname{cosech}^2 \kappa h) \cos^2 \theta d\theta, \quad (35)$$

with  $\kappa$  given in terms of  $\theta$  by (31)

This may be compared with (17) for the similar wave pattern in deep water

For a horizontal doublet  $M$  at depth  $f$  in water of depth  $h$ , an expression for the complete surface elevation can be derived from results given previously\* We have

$$\zeta = \frac{M}{\pi c} \int_{-\pi}^{\pi} d\theta \int_0^{\infty} \frac{\cosh \kappa(h-f) e^{i\kappa(x \cos \theta + y \sin \theta)}}{\cosh \kappa h (\kappa - \kappa_0 \sec^2 \theta \tanh \kappa h + i\mu \sec \theta)} \kappa^2 d\kappa, \quad (36)$$

where we take the limiting value of the real part for  $\mu \rightarrow 0$

From this we may easily deduce the free wave pattern to which the disturbance approximates at a great distance in the rear It is given by

$$\zeta = \frac{4\kappa_0 M}{c} \int_{-\pi}^{\pi} \frac{\cosh \kappa(h-f) \tanh^2 \kappa h \sec^4 \theta}{\cosh \kappa h (1 - \kappa_0 h \sec^2 \theta \operatorname{sech}^2 \kappa h)} \sin \{\kappa(x' \cos \theta + y \sin \theta)\} d\theta \quad (37)$$

From (35) this gives

$$R = 16\pi \rho \kappa_0 M^2 \int_0^{1\pi} \frac{\kappa^3 \cos \theta \cosh^2 \kappa(h-f)}{\cosh^2 \kappa h (1 - \kappa_0 h \sec^2 \theta \operatorname{sech}^2 \kappa h)} d\theta \quad (38)$$

It will be found that this agrees with the result obtained by a different method in the paper just quoted, when the previous expression is corrected for an obvious slip, in formula (37) of that paper 32 should be replaced by 16 and  $\tanh \kappa h (1 + \tanh \kappa h)$  by  $(1 + \tanh \kappa h)^2$

\* 'Proc Roy Soc., A, vol 118, p 32 (1928)

When  $\kappa_0 h < 1$ , that is  $c^2 > gh$ , the equation (31) for  $\kappa$  has a real root only for a more limited range of values of  $\theta$ , the lower limit being  $\theta_0 = \cos^{-1} \sqrt{(\kappa_0 h)}$  instead of zero. It is readily seen that the expression for  $R$  will be as in (38) with  $\theta_0$  as the lower limit of the integral

### Summary

An examination is made of the transfer of energy in a free wave pattern, and expressions for wave resistance are deduced. These are applied to certain cases both for deep water and for water of finite depth.

## Structures of the Metallic Carbonyl and Nitrosyl Compounds

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Our knowledge of the compounds containing carbonyl and nitrosyl (NO) groups attached to metallic atoms has been greatly extended in recent years, largely through the work of Hieber and Manchot and their collaborators, but the structures which have been suggested, especially for the nitrosyl derivatives, are not in every way satisfactory. In this paper an attempt is made to establish the structures of these compounds on a firmer basis.

### 1 Carbonyl Compounds

The nature of the linkage formed between carbon monoxide and a metallic atom is now well established. In its complexes the CO group occupies one co-ordination place as donor—it provides two electrons to form a single link—as is shown, for example, by the ferrous pentacyano-compounds of the general formula  $M_2[Fe(CN)_5X]^*$  where X may be CO,  $H_2O$ ,  $NH_3$ , pyridine, etc. This formation of a single link is in agreement with the Langmuir formula for carbon monoxide,<sup>†</sup>



which has been shown<sup>§</sup> to be supported by the electrical dipole moment, the

\* Gmelin-Kraut, "Handbuch d. anorg. Chem.," 'Iron,' B, pp 740, 1046, 1066, etc.

† 'J. Amer. Chem. Soc.,' vol 41, p. 1543 (1919).

‡ In the structural formulae given above, the charges assigned to the atoms are those which they would have if the electrons were equally shared. For the purpose of this paper the symbol  $\bar{\text{A}} - \dot{\text{B}}$  for the co-ordinate link is more convenient than  $\text{A} \leftarrow \text{B}$ , which means the same thing.

§ See Hammick, New, Sidgwick and Sutton, 'J. Chem. Soc.,' p 1878 (1930), Sidgwick, 'Chem. Rev.,' vol 9, p 77 (1931).

interatomic distance, the heat of formation, the force constant and the parachor. The CO molecule is obviously capable of co-ordinating as donor with another atom, either through the carbon or the oxygen. It is more probable that the link is formed through the carbon as in II, since (1) the 4-covalent condition is the normal state of the carbon atom, whereas it only rarely (e.g., in "basic" beryllium acetate  $\text{Be}_4\text{O}(\text{O}-\text{CO}-\text{CH}_3)_6$ ) occurs with oxygen, and (2) similar types of co-ordination compounds are formed by the isocyanides, for example with the cyanides of the metals Co, Ni, Cu, Ag, Cd, and with other salts, as in  $[\text{Pt}(\text{CH}_3\text{NC})_4]\text{PtCl}$  and  $[\text{Pt}(\text{CH}_3\text{NC})_2\text{Cl}_2]$ \*. Here the link must be formed through the carbon (which is in precisely the same state as in carbon monoxide), since the nitrogen has no unshared electrons. The arguments that follow would not, however, be affected if we supposed the carbonyl group to be attached through the oxygen.

This structure (II above), involving a triple link between the carbon and the oxygen, is strongly supported by the observation of Sutton and Bentley† that the electrical dipole moment of nickel carbonyl is zero. Unless the  $\text{M}-\text{C}-\text{O}$  group were linear, the molecule must have a moment owing to the rotation of the groups, as occurs with methyl and ethyl orthocarbonic esters  $\text{C}(\text{O}-\text{CO}-\text{R})_4$ . The fact that a CO group can replace an  $\text{H}_2\text{O}$  or  $\text{NH}_3$  in a complex shows that the link to the metal is single, and this is only compatible with a linear grouping of the atoms  $\text{M}-\text{C}-\text{O}$  if the  $\text{C}-\text{O}$  link is triple. The Raman spectrum of nickel carbonyl confirms the presence in it of a triple link of carbon to oxygen ‡.

We may therefore accept the structure  $\bar{\text{M}}-\text{C} \equiv \overset{+}{\text{O}}$  as established. Now five monometallic carbonyls (\*e., containing only one metallic atom in the molecule) are known

	$\text{Cr}(\text{CO})_6$ §	$\text{Fe}(\text{CO})_5$	$\text{Ni}(\text{CO})_4$	$\text{Mo}(\text{CO})_6$ ¶	$\text{W}(\text{CO})_6$ **
E A N	Cr 36	Fe 36	Ni 36	Mo 54	W 86

\* Hofmann and Bugge, 'Ber. deutsch. chem. Ges.', vol. 40, p. 1774 (1907), Tschuganov and Tsearu, *ibid.*, vol. 47, p. 570 (1914).

† 'Nature', vol. 130, p. 314 (1932), Sutton, New and Bentley, 'J. Chem. Soc.', p. 652, (1933).

‡ Anderson, 'Nature', vol. 130, p. 1002 (1932).

§ Job and Cassal, 'Bull. Soc. Chim.', vol. 41, p. 1041 (1927).

|| Mond and Langer, 'J. Chem. Soc.', vol. 59, p. 1091 (1931), Gmelin Kraut, "Handbuch d. anorg. Chem.," 'Iron,' B, p. 486.

¶ Mond, Hirts and Cowlap, 'J. Chem. Soc.', vol. 97, p. 798 (1910), Mond and Wallis, *ibid.*, vol. 121, p. 34 (1922). The analyses varied between  $\text{Mo}(\text{CO})_6$  and  $\text{Mo}(\text{CO})_5$ , but the analogy of the chromium and tungsten compounds fixes the composition.

\*\* Job and Rouvillois, 'C. R. Acad. Sci. Paris,' vol. 187, p. 564 (1922).

It is generally recognized that in all these the effective atomic number (E A N the total number of electrons, shared and unshared, in the molecule, in these compounds the atomic number increased by 2 for every CO) of the metallic atom is that of the next following inert gas, as is shown above\*. This fact, which explains why monometallic carbonyls are not formed by elements of odd atomic number, is very remarkable. As a rule it is found that the stability of a molecule depends, not on the total number of electrons in the E A N, but separately on the number in the core and that in the valency group. In these substances, however (which are all derivatives of the transitional elements of Periodic Groups VI, VII and VIII), the determining factor seems to be the total number, irrespective of how it is divided between the shared and the unshared electrons, thus the numbers are (the shared electrons are underlined) —

Cr(CO)	Cr =	2, 8, 14, <u>12</u> = 36
Fe(CO)	Fe =	2, 8, 16, <u>10</u> = 36
Ni(CO)	Ni =	2, 8, 18, <u>8</u> = 36
Mo(CO)	Mo =	2, 8, 18, 14, <u>12</u> = 54
W(CO)	W =	2, 8, 18, 32, 14, <u>12</u> = 86

It does not, however, seem to have been noticed that there is a further remarkable regularity in the composition of the carbonyls, which extends to those that contain more than one atom of metal in the molecule (the poly-metallic carbonyls). If we calculate the average E A N of the metallic atoms, by adding 2 to the atomic numbers for every CO, then the difference between the result and the atomic number of the next inert gas is always found to be one less than the number of metallic atoms in the molecule. If we write the compound  $M_n(CO)_y$ , and if  $m$  is the atomic number of  $M$ , and  $G$  that of the next inert gas, then the equation

$$G - \frac{x(m + 2y)}{x} = x - 1 \quad (1)$$

is always found to be true. Where  $x = 1$  (monometallic carbonyls) we have seen that this holds throughout. The other known pure carbonyls (containing only metallic atoms and CO groups) are the following, the molecular

\* This was given by Langmuir ('Science,' vol. 54, p. 65 (1921)) as the cause of the stability of iron and nickel carbonyls, and also accounts for the diamagnetism of these compounds, Oxley, 'Proc Camb Phil Soc,' vol 16, p. 102 (1911)

weights of all are known, and they are all derivatives of iron or cobalt, so that G always equals 36

$$\text{Fe}_2(\text{CO})_9^{\dagger} \quad \frac{2 \times 26 + 2 \times 9}{2} = 35 \quad x = 2$$

$$\text{Fe}_3(\text{CO})_{12}^{\dagger} \quad \frac{3 \times 26 + 2 \times 12}{3} = 34 \quad x = 3$$

$$\text{Co}_2(\text{CO})_8^{\dagger} \quad \frac{2 \times 27 + 2 \times 8}{2} = 35 \quad x = 2$$

$$\text{Co}_4(\text{CO})_{12}^{\S} \quad \frac{4 \times 27 + 2 \times 12}{4} = 33 \quad x = 4$$

This equation, with the appropriate modifications for other groups in the molecule, can be shown to hold also for the great majority of metallic derivatives containing carbonyl groups, and, if we accept the view of the nitrosyl group proposed in the second part of this paper, for the majority of the nitrosyl derivatives as well. We may fairly assume that in these polymetallic compounds the rule still applies that the E.A.N. is that of the next inert gas|| and that the necessary increase in the number of available electrons is secured by further co-ordination, which holds these atoms together. It is very improbable that the metallic atoms are united directly, as this form of linkage is practically unknown except with mercury, but it is obvious that the CO group is capable of forming a second co-ordinate link through the oxygen as donor, as in  $\text{M} \leftarrow \text{C} \equiv \text{O} \rightarrow \text{M}$  or  $\bar{\text{M}} - \overset{+}{\text{C}} \equiv \overset{+}{\text{O}} - \bar{\text{M}}$ . Since equation (1) holds for these compounds, it follows that when the number of M atoms in the molecule is 2, 3 and 4, the number of extra electrons needed in the molecule to make up the E.A.N. of every M to that of the next inert gas is 2, 6 and 12, and hence the number of new co-ordinate links required is 1, 3, and 6 respectively. This implies that every metallic atom in the molecule is joined to every other

\* Mond and Langer, 'J. Chem. Soc.' vol. 59, p. 1092 (1891), Dewar and Jones, 'Proc. Roy. Soc., A,' vol. 76, p. 573 (1905), Speyer and Wolf, 'Ber. deuts. chem. Ges.,' vol. 60, p. 1424 (1927).

† Mond, Hirtz and Cowap, 'J. Chem. Soc.,' vol. 97, p. 798 (1910).

‡ Dewar and Jones, 'Proc. Roy. Soc., A,' vol. 79, p. 75 (1907), Freundlich and Cuy, 'Ber. deuts. chem. Ges.,' vol. 56, p. 2265 (1923).

§ Hieber, Mühlbauer and Ehmann, 'Ber. deuts. chem. Ges.,' vol. 65, p. 1090 (1932).

|| This is supported by the fact that  $\text{Fe}_2(\text{CO})_9$  and  $\text{Fe}_3(\text{CO})_{12}$  are diamagnetic, like  $\text{Fe}(\text{CO})_5$  and  $\text{Ni}(\text{CO})_4$ , Freundlich, 'Ber. deuts. chem. Ges.,' vol. 56, p. 2284 (1923), Berkmann and Zocher, 'Z. phys. Chem.,' vol. 124, p. 318 (1926).



through a link of this kind. For two such atoms this is secured by joining them through a single CO, for three, by placing them at the angular points of a triangle with a CO in each side, and for four by placing them at those of a tetrahedron, with a CO group on each edge.

These curious relations may be regarded in two ways: either (a) they depend on the validity of equation (1), or (b) they are connected with the linear, triangular, and tetrahedral structures. For  $x = 2, 3$ , or  $4$ , either hypothesis gives the same result, but for higher values of  $x$  they differ. If equation (1) is the determining factor, then since that involves a link between every metallic atom and every other, and since for physical reasons these links must obviously be of the same length, no value of  $x$  greater than  $4$  is possible, for we cannot have more than four points equidistant from one another in space. As a fact no carbonyls with more than four metallic atoms in the molecules are known, and this may be the reason, or it may only be an accident that they have not yet been discovered. If the real principle is the adoption of a series of simple geometrical forms (together with the attainment of the inert gas number), then we might expect the tetrahedron to be followed by an octahedron, and that by a cube. For an octahedron, with a CO on each edge, the formulae which would give each M an inert gas number are  $\text{Fe}_6(\text{CO})_{12}$ ,  $\text{Co}_8(\text{CO})_{12}$ , and  $\text{Ni}_4(\text{CO})_{12}$ . These do not satisfy equation (1), which would require  $x$  to be  $5$ , not  $6$ . A cube, with  $8$  points and  $12$  sides, would give the same result as a tetrahedron ( $4$  and  $6$ ) and so is not likely to occur.

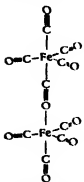
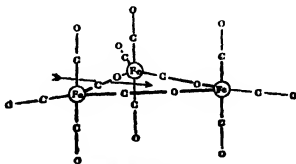
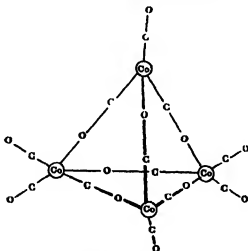
The structures assigned on this hypothesis to  $\text{Fe}_2(\text{CO})_9$ ,  $\text{Fe}_3(\text{CO})_{12}$ , and  $\text{Co}_4(\text{CO})_{12}$ ,  $\text{Fe}_4(\text{CO})_{14}$ , which we should expect to exist, have not been obtained as given in figs. 1 to 3.

The crystal structures of the first two of these structures have been examined by Brill\*. We are much indebted to Mr. H. M. Powell for the following note on Brill's results:—

With  $\text{Fe}_2(\text{CO})_9$  Brill finds that the substance crystallizes in the hexagonal system, and concludes that the symmetry of the molecule is either  $C_2$ ,  $C_3$ ,  $C_{2h}$ ,  $D_3$ , or  $D_{3h}$ , but he does not determine the atomic positions. The structure he suggests is different from that given in fig. 1, and has three layers of three CO's each, with the two iron atoms, which must both lie on a threefold symmetry axis, between them. But in a note (p. 89) he says "It is possible to bring some of the CO's into onefold positions." This leads to crowding of the threefold axis, and is chemically

\*  $\text{Fe}_2(\text{CO})_9$ , 'Z. Kristallog.' vol. 65, p. 80 (1927),  $\text{Fe}_3(\text{CO})_{12}$ , *ibid.*, vol. 77, p. 36 (1931).

not very probable." The proposed formula I has the symmetry  $C_{2v}$ , or, if one group of three CO's is rotated relative to the other through an arbitrary angle,  $C_2$  it could therefore be put into some of the possible space groups found by Brill. We have no independent evidence of the disposition in space of five covalencies (which cannot in any event be

FIG 1.— $Fe_3(CO)_9$ .FIG 2.— $Fe_3(CO)_{12}$ .FIG 3.— $Co_4(CO)_{12}$ .

symmetrically equivalent in three dimensions), but that given in fig 1 is at least as probable as any. Moreover, the dimensions required by the formula agree with Brill's observations. The long vertical axis is  $O=C-Fe-C \equiv O-Fe-C \equiv O$ , if we may assume the two valencies of the iron to be at  $180^\circ$  this must be linear, as the X-ray results show it to be, if we take the lengths of the links to be  $C \equiv O$

1 14 C — Fe 0 77 + 1 27 = 2 04 O — Fe 0 70 + 1 27 = 1 97,\* the distance between the centres of the terminal oxygen atoms should be 11 5 Å. The whole height of the cell according to Brill is 15 8 Å, which means that the distance between the centres of two neighbouring oxygens in different molecules is 4 3 Å. This is about the usual value, in graphite two carbon atoms in different sheets (*i.e.*, molecules) are 3 4 Å apart, in benzene hexachloride two such chlorines are 3 74 Å apart†. Thus there is no undue crowding on the threefold axis, and the structure, fig 1, proposed for  $\text{Fe}_2(\text{CO})_9$  in no way conflicts with the X-ray evidence.

With  $\text{Fe}_3(\text{CO})_{12}$  the X-ray evidence is less easy to interpret. The molecule has symmetry  $C_3$ . Formula II, fig 2, can be given symmetry very near to  $C_3$  if we distort the triangle slightly in the direction of the arrow shown in the figure, which then represents the twofold axis. This is on the assumption that a carbon and an oxygen atom are indistinguishable, which is probable since on the proposed structure each has the same number of electrons. The formula which Brill prefers requires at least as much distortion to make it fit the observations.

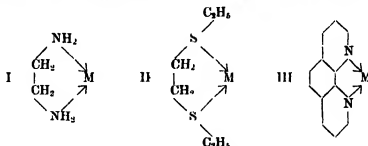
It might be objected to these formulae that they involve too great a strain in the ring, the chain of atoms  $\text{M} - \text{C} \equiv \text{O} - \text{Fe}$  being in its unstrained condition linear. But we may assume that the natural angle between the two nearest valencies of a 5-covalent atom does not exceed  $90^\circ$ , as in fig 1. Hence in the triangle, fig 2, the total strain is  $3(90^\circ - 60^\circ) = 90^\circ$ , and this is divided among 9 atoms, giving only  $10^\circ$  per atom, which is to be compared with the strain of  $50^\circ$  per atom in cyclopropane, or  $20^\circ$  per atom in cyclobutane.

In addition to the pure carbonyls, there are a large number of mixed derivatives which contain CO groups, and of these the majority, though not all, when their E.A.N.'s are calculated in the usual way, give the values required by equation (1), they must therefore be supposed to have similar structures. Thus we can have one or more of the CO groups replaced by an equal number of other donor molecules such as ammonia or pyridine (*py*), or two CO groups by a chelate molecule (*i.e.*, one containing two donor atoms in positions suitable for forming a ring) such as ethylene diamine (*en*), the di-ethyl ether of dithioglycol (*th*) (II), or *o*-phenanthroline (*phen*) (III).

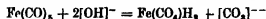
\* These distances are taken from Sidgwick, "The Covalent Link," pp. 85, 88 (1933).

† Hendricks and Billike, 'J. Amer. Chem. Soc.', vol. 48, p. 3007 (1926).

Examples are  $\text{Fe}(\text{CO})_4(\text{py})\ddagger$ ,  $\text{Fe}(\text{CO})_3(\text{py})_2\ddagger\ddagger$ ,  $-(\text{NH}_3)_2\S$ ,  $\text{Fe}(\text{CO})_2(\text{en})\|\text{**}$ ,  $-(\text{phth})\dagger\dagger$ ,  $\text{Ni}(\text{CO})_3(\text{phth})\ddagger\ddagger$ , and among the polymetallic compounds  $\text{Fe}_2(\text{CO})_9(\text{en})_2^*\text{**}$ ,  $-(\text{phth})\dagger\dagger$  and  $\text{Ni}_2(\text{CO})_8(\text{py})_2\ddagger\ddagger$ . In all these the E A N remains 36, and the covalency that of the simpler carbonyls (5 for iron, 4 for



nickel) In another series we find the E A N preserved, but the covalency increased for iron from 5 to 6. The most remarkable of these is the volatile hydrogen compound  $\text{H}_2\text{Fe}(\text{CO})_4\S\S$ . This is a volatile unstable yellow liquid melting about  $-68^\circ\text{C}$ , which is formed by the action of alkalis on iron pentacarbonyl



It is obvious from its volatility that it is covalent, and it gives a series of derivatives in which the hydrogens are replaced by halogens, as in  $\text{Fe}(\text{CO})_4\text{Cl}_2$ ,  $-\text{Br}_2$ ,  $-\text{I}_2\|\|$  or by mercury, as in  $\text{Fe}(\text{CO})_4\text{Hg}$ ,  $\text{Fe}(\text{CO})_4(\text{HgCl})_2$ ,  $-(\text{HgBr})_2$ ,  $-(\text{HgI})_2\|\|$  or they can be replaced by metals with the formation of salts, and the reduction of the covalency to 5 or 4, as in  $\text{Fe}(\text{CO})_4\text{H}[\text{Na}]$ ,  $\text{Fe}(\text{CO})_4\text{H}_2[\text{Ba}]$ ,  $\text{Fe}(\text{CO})_4[\text{Cd}]\P\P$ . Similar derivatives are known with several metallic atoms in the molecule, and these still satisfy equation (1), examples are  $\text{Fe}_2(\text{CO})_7\text{Br}_2\|\|$  and  $\text{Fe}_3(\text{CO})_9\text{Br}_6$  (molecular weight determined)  $\P$ . We also find compounds containing both halogens and co-ordinated molecules, such as  $\text{Fe}(\text{CO})_2\text{Cl}_2(\text{phth})$ ,  $-\text{Br}_2(\text{phth})$ ,  $\text{Fe}(\text{CO})_2\text{I}_2(\text{th})$ ,  $-(\text{py})_2\P$ .

\* Hieber and Sonneckalb, 'Ber. deut. chem. Ges.', vol. 61, p. 558 (1928).

† Hieber and Bader, *ibid.*, p. 1717 (1928).

‡ Hieber and Sonneckalb, *ibid.*, p. 2421.

§ Hieber, 'Sitzber. heidelb. Akad. Wiss.', vol. 3, p. 4 (1929).

|| Hieber and Becker, 'Ber. deut. chem. Ges.', vol. 63, p. 1405 (1930).

¶ Hieber, 'Z. anorg. Chem.', vol. 201, p. 329 (1931).

\*\* Hieber and Leutert, 'Ber. deut. chem. Ges.', vol. 64, p. 2832 (1931).

†† Hieber and Mühlbauer, *ibid.*, vol. 65, p. 1082 (1932).

‡‡ Hieber, Mühlbauer and Ehmman, *ibid.*, p. 1090.

§§ Hieber and Leutert, 'Naturwiss.', vol. 19, p. 360 (1931).

||| Gmelin Kraut, 'Handbuch d. anorg. Chem.', 'Iron,' B, pp. 499-502.

¶¶ Feigl and Krumholz, 'Z. anorg. Chem.', vol. 215, p. 242 (1933).

Some examples of the working out of the E.A.N. and of the size of the valency group may be given —

$\text{Fe}(\text{CO})_5(\text{en})$	$26 + 2 \times 3 + 4$	$= 36$	2, 8, 16, <u>10</u> ,
$\text{H}_2\text{Fe}(\text{CO})_4$	$26 + 2 + 2 \times 4$	$= 36$	2, 8, 14, <u>12</u> ,
$\text{Fe}(\text{CO})_4\text{H}[\text{Na}]$	$26 + 2 \times 4 + 1 + 1$	$= 36$	2, 8, 16, <u>10</u> ,
$\text{Fe}_2(\text{CO})_8(\text{en})_2$	$\frac{2 \times 26 + 2 \times 5 + 2 \times 4}{2}$	$= 35$	$x = 2$ ,
$\text{Fe}_3(\text{CO})_9\text{Br}_4$	$\frac{3 \times 26 + 2 \times 9 + 6}{3}$	$= 34$	$x = 3$

The rule is obeyed equally by other series of compounds in which CO replaces water or ammonia in recognized complexes, as in the pentacyano-compounds  $\text{M}_3[\text{Fe}(\text{CH})_3(\text{CO})]$  already mentioned. It is clear that this rule expresses the condition of stability of metallic atoms of the transitional elements attached to carbonyl groups.

## II Nitrosyl Compounds

The nitrosyl compounds have one or more NO groups attached to a metallic atom. In their preparation and properties they show a close analogy to the carbonyls. They are commonly formed by the same metals, they are usually prepared by the direct action of nitric oxide, as the carbonyls are by that of carbon monoxide, and often these two gases can drive one another out of their respective compounds\*. We may therefore expect to find that the nitrosyls and the carbonyls have analogous structures.

The suggestion that the metallic nitrosyls are derivatives of hyponitrous acid

$$\begin{array}{c} \text{N}-\text{OH} \\ || \\ \text{N}-\text{OH} \end{array}$$

has been sufficiently refuted by Manchot†. They are formed from nitric oxide and not from hyponitrous acid, except under conditions where the latter forms nitric oxide, they readily liberate nitric oxide (as the carbonyls liberate carbon monoxide) with acids, whereas the hyponitrites with acids give nitrous oxide. Also this assumption would require us to double the formulae of many of them, such as the nitroprussides and generally those which have

\* Manchot and Schmidt, 'Z. anorg. Chem.', vol. 216, p. 99 (1933).

† Manchot and Davidson, 'Ber. deuts. chem. Ges.', vol. 62, p. 684 (1929). See also Manchot and Schmidt, 'Z. anorg. Chem.', vol. 216, p. 103 (1933). It is not impossible that metallic complexes derived from hyponitrous acid may occur, but they are not found among the normal nitrosyls.

one NO to one metallic atom, there is no evidence in favour of this, and much against it

We may therefore conclude that each NO group is separately attached to the metal, and the first question is what is the nature of the attachment. It can be shown that this is a single co-ordinate link  $M \leftarrow$  by the fact that it is possible to replace a CO by an NO in a series of complexes of the co-ordination number 6. Examples of these are given in Table I

Table I

$K_2[Mn(CN)_5(NO)]^*$ $K_2[Fe(CN)_5(NO)]$  $K_2[Ru(CN)_5(NO)]^\dagger$ $K_2[RuCl_4(NO)]^\ddagger$ $[Ru(NH_3)_4Cl(NO)] Br_2$  $K_2[OsCl_4(NO)]  $	$K_2[Fe(CN)_5(CO)]$ $K_2[Fe(CN)_5(H_2O)]$	$K_2[Mn(CN)_6]$ $K_2[Fe(CN)_6]$  $K_2[Ru(CN)_6]$   $K_2[OsCl_6]$ $K_2[Os(CN)_6]$
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\* Manchot and Schmidt, 'Ber. deut. chem. Ges.', vol. 59, p. 2360 (1926)

† Manchot and Düsing, *ibid.*, vol. 63, p. 1226 (1930)

‡ Joly, 'O. R. Acad. Sci. Paris', vol. 107, p. 994 (1888), Howe, 'J. Amer. Chem. Soc.', vol. 16, p. 388 (1894), Manchot and Schmidt, 'Z. anorg. Chem.', vol. 216, p. 99 (1933)

§ Rosenbohm, 'Z. phys. Chem.', vol. 93, p. 693 (1919)

|| Wintrobort, 'Ann. Chem. Phys.', vol. 28, p. 15 (1903)

Of these compounds the nitroprussides  $M_2[Fe(CN)_5(NO)]$  are the most familiar, and in this anion, as we have seen, the NO can be replaced not only by CO, but also by  $H_2O$ ,  $NH_3$ , and other molecules, the replacement is always accompanied by a fall in the electrovalency of the anion, which will be discussed later. The suggestion that the NO is doubly linked to the metal cannot be entertained. The co-ordination number 6 (group of 12 shared electrons) is universal in complexes of the pentacyano-type, and is, as Werner showed, the commonest of all complex forms, a co-ordination number 7 is almost unknown throughout chemistry, and has never been observed with these transitional metals. Moreover, for iron and manganese a co-ordination number greater than 6 is not only unknown, but is impossible according to the covalency rule\*. We may therefore take it as proved that the metal is joined to the NO by a single co-ordinate link. That the attachment to the metal is through the nitrogen is shown by the fact that the nitroso-group in these compounds can be oxidized to a nitro-group.

If we accept these conclusions, there are three structures possible for the  $M-N-O$  group

\* Sidgwick, "Electronic Theory of Valency," p. 152 (1927).

(1) It may contain a true nitroso-group  $M-N=O$ , as in the organic nitroso-compounds. This view cannot be maintained, the nitrosyls are devoid of the characteristic tendency of the nitroso-compounds to polymerize with loss (or change) of colour, also this structure would involve the assumption of valencies for the metallic atoms for which there is no parallel elsewhere, such as the quadrivalency of iron in the nitroprussides. Moreover, Werner and Karrer\* have obtained a complex salt which has the characteristics of such a true nitroso-derivative. This is  $[Co(NH_3)_5(NO)]X_2$ , a series of salts were prepared. They occur in a black monomolecular form which changes to a red dimeric form. These belong to quite a different type from the nitrosyls, regarded as nitroso-compounds they correspond to  $[Co(NH_3)_5Cl]Cl_2$ , and the cobalt is trivalent, as it normally is in its 6-co-ordination compounds.

(2) It might be supposed that they are formed by simple co-ordination, by the sharing of a lone pair of electrons of the nitrogen in the nitric oxide molecule with the metal, without any further change occurring. Then the mysterious link—presumably of 5 electrons—which exists in nitric oxide must be maintained in the nitrosyls. This link, however, is, so far as we know, except in nitric oxide and perhaps in a few of its organic derivatives, too unstable to exist, and it may be expected to pass over in the complex into some more normal form. Moreover on this hypothesis that the  $N-O$  link is the same in the complex as in nitric oxide (which is the one most generally accepted for the nitroprussides), the replacement of  $CO$  by  $NO$  should involve no other change in the molecule, whereas in fact it is always accompanied by a change in the electrovalency, as from  $K_3[Fe(CN)_5(CO)]$  to  $K_2[Fe(CN)_5(NO)]$  (see Table I). This second hypothesis is quite unable to account for such changes.

(3) The close similarity between nitric oxide and carbon monoxide, and between the nitrosyls and the carbonyls, suggests that the structures of the latter are similar, and that there is a triple link between the nitrogen and the oxygen giving  $M-N \equiv O$ , corresponding to  $M-C \equiv O$ . But nitric oxide has one more electron than carbon monoxide (balanced of course by an extra positive charge on the nitrogen nucleus), this electron, if it does not remain as part of an "odd-electron"  $N-O$  link, must go somewhere. We have evidence of this tendency of the  $NO$  link to lose an electron in the existence of positive  $[NO]^+$  ion, which has been established by Hantzsch and Berger†. They have shown that nitrosyl perchlorate  $[NO]ClO_4$  is anhydrous, and gives a conducting solution in nitromethane, and that the so-called "nitrosulphonic

\* 'Helv. Chem. Acta,' vol 1, p 54 (1918).

† 'Z. anorg. Chem.', vol 190, p. 321 (1930).

acid" which is usually written  $\text{NO}_2\text{SO}_3\text{H}$ , is dissociated in concentrated sulphuric acid, and is no doubt nitrosyl hydrogen sulphate  $[\text{NO}]\text{SO}_3\text{H}$ . Another salt of this type is nitrosyl fluoborate  $[\text{NO}]\text{BF}_4$ , obtained\* by treating concentrated aqueous fluoboric acid with oxides of nitrogen, which has been shown† to have this composition and to be free from water. It is thus evident that a cation  $[\text{NO}]^+$  can exist. This forms with  $[\text{CN}]$  and  $\text{CO}$  a series of diatomic molecules with identical electronic arrangements, as shown in Table II. The positions of the electrical charges, on the simplifying assumption that the electrons are shared equally, are given in the second line of symbols, every atom in these molecules has 2 unshared and 6 shared electrons, equivalent to a total of 5, and this gives the residual charges as  $\text{C} - 1$ ,  $\text{N}$  zero,  $\text{O} + 1$ . If we suppose a combination of the neutral groups  $\text{CN}$ ,  $\text{CO}$ ,  $\text{NO}$  with a metallic atom  $\text{M}$ , we get the structures given in the table for  $\text{M}-\text{C}-\text{N}$ , etc., the  $\text{CN}$  group contributes one electron to the  $\text{E A N}$  of the metal (normal covalency), the  $\text{CO}$  two (co-ordinate link), while the  $\text{NO}$ , in addition to the 2 for the co-ordinate link, adds a third, by the complete transference of an electron from the  $\text{NO}$  to the  $\text{M}$ , and thus contributes three. These charges diminish the electrovalency of the complex by 1 and 2 respectively, as is shown in the formulæ of the pentacyano-compounds at the foot of the table.

Table II

$[\text{C N}]^-$	$[\text{C O}]$	$[\text{N O}]^+$
$\bar{\text{C}} \quad \text{N}$	$\bar{\text{C}} \quad \bar{\text{O}}$	$\text{N} - \bar{\text{O}}$
$\text{M C N}$	$\text{M C O}$	$\text{M N O}$
$\text{M} - \text{C} \equiv \text{N}$	$\text{M} - \text{C} \equiv \bar{\text{O}}$	$\text{M} - \text{N} \equiv \bar{\text{O}}$
$\text{K}_4[\text{Fe}(\text{CN})_6]$	$\text{K}_5[\text{Fe}(\text{CN})_5(\text{CO})]$	$\text{K}_6[\text{Fe}(\text{CN})_5(\text{NO})]$

If we make this assumption that the  $\text{NO}$  group, though it is only attached to the metal by a single link of two shared electrons, yet contributes a further electron, and counts as 3 towards the  $\text{E A N}$  of  $\text{M}$ , the whole series of nitrosyl compounds given in the left-hand column of Table I is brought into line with the other complexes of these metals, in the other two columns ‡

\* Wilke Dörfurt and Balz, *ibid.*, vol 159, p 197 (1927)

† 'Z anorg Chem,' vol 190, p 321 (1930)

‡ Reiff, 'Z anorg Chem,' vol 202, p 376 (1931) suggests that this contribution of three electrons made by the  $\text{NO}$  to the  $\text{E A N}$  of the metallic atom explains the existence and the diamagnetism of  $\text{Co}(\text{CO})_5(\text{NO})$  (see below), but he does not seriously discuss the nature of the linkage



Thus the nitroprussides are  $M_3[Fe(CN)_5(NO)]$  while the corresponding carbonyl compounds are  $M_3[Fe(CN)_5(CO)]$ , if we allow for the extra electron going to the iron in the former, this makes the nitroprussides, like their carbonyl analogues, ferrous and not ferric compounds. Now among all the known compounds of the type  $M_n[Fe(CN)_5X]$ , where X may be CN,  $NO_2$ ,  $AsO_3$ ,  $SO_3$ ,  $H_2O$ ,  $NH_3$ , or CO,\* it is always found that the ferrous are more stable than the ferric, the latter when they occur have ferrous analogues, but the reverse is not true, in particular, there is no ferric analogue of  $M_3[Fe(CN)_5(CO)]$ . It is therefore definitely in favour of this view that it represents the nitroprussides as ferrous compounds, the anomaly of a stable ferric pentacyano-compound with no ferrous equivalent is thereby removed. In the same way the manganese in the compound in Table I has the same valency as in the colourless cyanide  $K_2[Mn(CN)_6]$ , and the ruthenium compound as in the only stable cyanide of ruthenium  $K_4[Ru(CN)_6]$ . Further, the E A N of the elements in Table I is Mn, Fe 36 Ru 24 Os 86, that is, the nitrosyl like the carbonyl derivatives follow the inert gas rule.

Another way of retaining the E A N when a CO is replaced by an NO is to replace at the same time the central atom by an atom with an atomic number less by 1. This is illustrated by the remarkable series of volatile compounds analogous to nickel carbonyl

	$Ni(CO)_4$ °C	$Co(CO)_5(NO) \dagger$ °C	$Fe(CO)_5(NO)_2 \ddagger$ °C
B Pt	43	78.6	110
M Pt	-23	-1.1	+18

Hieber and Anderson who prepared the iron compound from iron enneacarbonyl  $Fe_9(CO)_9$ , draw attention to this parallelism, and recognize (as Reiff also does with respect to the cobalt compound) that the NO group contributes an extra electron, they point out that we might expect to get  $Mn(CO)(NO)_3$  and  $Cr(NO)_4$ , although the marked instability of the iron derivative make this rather improbable. They suppose, however, that the NO is attached by a double link (four shared electrons) to the metal, which would make the cobalt 5- and the iron 6-covalent. This is not only an unnecessary complication, but

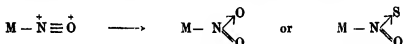
\* See Gmelin Kraut, "Handbuch d. anorg. Chem., 'Iron,' B, pp. 732-742.

† Mond and Wallis, 'J. Chem. Soc.', vol. 121, p. 34 (1922), Reiff, 'Z. anorg. Chem.', vol. 202, p. 375 (1931), Hieber and Anderson, *ibid.*, vol. 211, p. 132 (1933).

‡ Hieber and Anderson, *loc. cit.*, and *ibid.*, vol. 208, p. 238 (1932), the boiling point of the cobalt compound is given by mistake as 48.6°.

as we have seen it cannot be applied to the manganese and iron nitrosyl compounds in Table I, because it would give them a covalency of 7, which is above their maximum. The marked rise in the boiling point as we go from the nickel to the iron compound is sufficiently accounted for by the considerable dipole moment which the  $\bar{M} - \overset{+}{N} \equiv \overset{+}{O}$  group must give to the molecule.

This method of formulation further explains the characteristic colour reaction of the nitroprussides with the sulphides. They do not react with hydrogen sulphide, but on addition of alkali or of an alkaline sulphide to the solution the deep purple colour of  $M_4[Fe(CN)_6(NOS)]$  at once appears, showing that the reaction is with the sulphide ion\*. In the same way hydroxyl ions convert the nitroprussides into nitro-compounds  $M_4[Fe(CN)_6(NO_2)]$ †. It is evident that the strongly positive  $\overset{+}{N} \equiv \overset{+}{O}$  group attracts the negative  $OH^-$  or  $S^-$  (or  $HS^-$ ) ions, with the production of a nitro or thionitro-group



This reaction involves the withdrawal of two electrons from the central atom, and hence, in order to maintain the E.A.N., an increase of the electrovalency by two, the products having four atoms of M instead of two in the molecule.

It is particularly to be observed that in all the nitrosyl compounds hitherto mentioned this method of formulation involves the inert gas rule being maintained. This rule is so nearly universal among the carbonyls that it is a strong support of our hypothesis that it involves its extension to the nitrosyls as well. For a molecule  $M_x(CO)_y(NO)_z$  we get the equation

$$G - \frac{xm + 2y + 3z}{x} = x - 1, \quad (2)$$

which expresses the composition of the carbonyl-nitrosyls of iron and cobalt given above. Many mixed nitrosyl derivatives comply with this equation, such as  $Fe(NO)_3(phth)$  and  $Co(CO)(NO)(phth)$ ‡. It is important to notice that equation (2) holds for nitrosyls containing more than one metallic atom.

\* Virgili, 'Z. analyt. Chem.', vol. 45, p. 409 (1906).

† Cambi and Szegő, 'Rend. Acc. Lincei,' vol. 5, p. 737 (1927), 'Gazz. Chim. Ital.', vol. 58, p. 71 (1928). This is usually formulated as a nitrito-compound  $M - O - N = O$ , but according to Werner, "Neuere Anschauungen," pp. 131, 339 (1913), the stable form when M is a transitional element is the true-nitro-form  $M - NO_2$ .

‡ Heber and Anderson, 'Z. anorg. Chem.', vol. 211, p. 132 (1933).

in the molecule. One example is  $\text{Fe}_2(\text{NO})_4(\text{py})_2$ ,\* here, since each pyridine molecule, like a CO, adds two electrons, we have

$$g - \frac{2 \times 26 + 3 \times 4 + 2 \times 3}{2} = 36 - 35 = 1$$

( $x = 2$ )

Others are the curious "red and black salts of Roussin," which have long been known†. Their formulæ are  $\text{M}_2[\text{Fe}_2(\text{NO})_4\text{S}_2]$  (red, unstable) and  $\text{M}[\text{Fe}_4(\text{NO})_7\text{S}_3]$  (black, stable). The black salts are obtained by the action of nitric oxide or nitrites on iron salts in the presence of sulphides, alkalies convert them into the red series, which are unstable and readily revert to the black. The molecular weights of the black salts have been confirmed by conductivity measurements‡. Those of the red series are known through their esters, which Hofmann and Wiedes§ prepared by the action of nitric oxide on a mixture of ferrous sulphate and ethyl mercaptan, they found their molecular weight in solution to correspond to the formula  $(\text{C}_2\text{H}_5)_2\text{Fe}_2(\text{NO})_4\text{S}_2$ . On the probable assumption that the sulphur is doubly linked to the metal (in the ion as  $\text{Fe} = \text{S}$ , contributing two electrons, in the ester as  $\text{Fe} = \text{S}^+ - \text{C}_2\text{H}_5$ , contributing three, and so compensating for the disappearance of the ionic charge) these compounds satisfy equation (2)

$$\begin{array}{ll} \text{M}_2[\text{Fe}_2(\text{NO})_4\text{S}_2] & \frac{2 \times 26 + 3 \times 4 + 2 \times 2 + 2}{2} = 35 \quad x = 2 \\ \text{Fe}_2(\text{NO})_4(\text{S} - \text{C}_2\text{H}_5)_2 & \frac{2 \times 26 + 3 \times 4 + 2 \times 3}{2} = 35 \quad x = 2 \\ \text{M}[\text{Fe}_4(\text{NO})_7\text{S}_3] & \frac{4 \times 26 + 3 \times 7 + 3 \times 2 + 1}{2} = 33 \quad x = 4 \end{array}$$

This implies for the black salts a tetrahedral structure, like that given for  $\text{Co}_4(\text{CO})_{12}$  in fig 3

Among the nitrosyl compounds described in the literature there are a number which do not comply with this rule. Some could be brought into line with it by assuming a sufficient degree of polymerization, and the substances in

\* Hieber and Anderson, *Z anorg. Chem.*, vol 211, p 132 (1933)

† Gmelin Kraut, *loc cit.*, 'Iron,' B, pp 471-477, Manchot and Lanckh, *Ber. deutsch. chem. Ges.*, vol 59, p 407 (1926), Manchot and Gall, *ibid.*, vol 60, p 2318 (1927), Manchot and Lehmann, *Liebigs Ann.*, vol 470, p 255 (1929)

‡ Marchlewski and Sachs, *Z anorg. Chem.*, vol 2, p 181 (1892), Bellucci and Carnevali, *Rend. Acc. Lincei*, vol 16, p 654 (1907)

§ *Z anorg. Chem.*, vol 9, p 301 (1895)

question are obviously polymerized, but we do not know to what extent. Some give support to our general theory of the structure of the nitrosyl group, though not to the particular rule. Such are the carbonyl and nitrosyl derivatives of copper. The molecular weights and hence the complete structures of these compounds are unknown, but it is clear that the stable forms contain one carbonyl\* or one nitrosyl† group to one copper atom. Now the carbonyl compounds are all derived from cuprous copper, as  $\text{Cu}(\text{CO})\text{Br}$ , and the nitrosyls from cupric copper, as  $\text{Cu}(\text{NO})\text{Cl}_2$  and  $\text{Cu}(\text{NO})\text{SO}_4$ . It is evident that the extra electron contributed by the NO balances the defect in the copper. Other substances, such as Manchot's iron tetranitrosyl  $(\text{Fe}(\text{NO})_4)_2$ ,‡ must obviously have some totally different structure. But it is submitted that the method of formulation which we have proposed accounts for the composition and behaviour of the great majority of nitrosyl compounds, and brings their structures into conformity with those of the very similar carbonyls.

### Summary

I *Carbonyl Compounds*—These contain the group  $\bar{\text{M}} - \text{C} \equiv \overset{+}{\text{O}}$ . For all carbonyls  $\text{M}_x(\text{CO})_y$ ,  $G - \frac{x(m+2y)}{x} = x - 1$  ( $m$  = atomic number of M, G of next inert gas)  $x = 1, 2, 3$ , or 4. When  $x = 1$ , M has effective atomic number (E.A.N.) of inert gas. Assuming this to be true when  $x > 1$ , M atoms must be held together thus  $\bar{\text{M}} - \overset{+}{\text{C}} \equiv \overset{+}{\text{O}} - \bar{\text{M}}$ , if the molecule contains 2M they must have one such link, if 3, they must be at the angles of a triangle with a CO in each side, if 4, at the points of a tetrahedron with a CO on each edge. The crystal structures of  $\text{Fe}_2(\text{CO})_9$  and  $\text{Fe}_3(\text{CO})_{12}$  are compatible with this. Nearly all carbonyl derivatives follow these rules, with necessary modifications for the other groups present.

II *Nitrosyl Compounds*—In  $\text{M} - \text{N} - \text{O}$  the nitrogen is attached by two shared electrons to M. But when NO replaces CO in a complex the negative electrovalency falls by 1.  $\text{M}_2[\text{Fe}(\text{CN})_5(\text{CO})]$ ,  $\text{M}_2[\text{Fe}(\text{CN})_5(\text{NO})]$ . This is explained if the structure is  $\text{M} - \text{N} = \text{O}$  or  $\bar{\text{M}} - \overset{+}{\text{N}} \equiv \overset{+}{\text{O}}$ , corresponding to  $\text{M} - \text{C} = \text{O}$  or  $\bar{\text{M}} - \text{C} \equiv \overset{+}{\text{O}}$ , and the NO contributes three electrons to the E.A.N. of M, as CO contributes two.

\* Wagner, 'Z. anorg. Chem.', vol. 196, p. 364 (1931).

† Manchot, 'Liebigs Ann.', vol. 375, p. 308 (1910), 'Ber. deutsch. chem. Ges.', vol. 47, p. 1601 (1914), Manchot and Linckh, *ibid.*, vol. 59, p. 407 (1926).

‡ Manchot and Enk, 'Liebigs Ann.', vol. 470, p. 275 (1929).

On this hypothesis the structures of the great majority of nitrosyl derivatives are shown to follow precisely the same rules as the carbonyls. The changes in electrovalency of the anions, and the similarity of the volatile compounds  $\text{Ni(CO)}_4$ ,  $\text{Co(CO)}_2(\text{NO})$ ,  $\text{Fe(CO)}_2(\text{NO})_2$  are explained, and the nitrosyls with several metallic atoms in the molecules are shown to obey an equation corresponding to that given above

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### *An Annual Perturbation in the Range of Tide*

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#### 1 *Introduction*

In the following paper the existence is established of an annual perturbation in the range of tide. The perturbation became evident during an intensive examination, by a method\* described in the paper, of the residual semi-diurnal tide at Liverpool. Direct analysis† of hourly heights for an annual perturbation in the principal constituent of semi-diurnal tide showed the perturbation to be consistent from year to year, and to exist generally in British Waters, and throughout the world at large. Analysis of high and low waters further confirmed the results. A study of the general distribution of the perturbation provided a number of interesting results, as also did a study of its relation with local meteorological conditions.

The paper concludes with a short discussion on possible causes of an annual perturbation in the range of tide.

#### 2 *Observed and Synthesized Tides*

It is a noteworthy feature of Doodson's method of analysis in that it provides for each species of tide (diurnal, semi-diurnal, etc.) two numerical quantities for each solar day. Thus the hourly heights are treated in the

\* The method, which is new, is described in detail in Appendix I. It is dependent upon Doodson's method of tidal analysis, and was first indicated in his paper on the "Analysis of Tidal Observations," 'Phil. Trans.' A, vol. 227, p. 223 (1928).

† See Appendix II.

first place as though the tide were solar and the semi-diurnal part of it expressible in the form

$$R \cos (30^\circ t - \epsilon) = P \cos 30^\circ t + Q \sin 30^\circ t,$$

where

$t$  = the time in solar hours from a fixed hour of the day,

$\epsilon/30$  = the time of high water in solar hours,

$$P = R \cos \epsilon \quad \text{and} \quad Q = R \sin \epsilon$$

The numerical processes of analysis give each day two quantities, related to  $P$  and  $Q$ , which are almost entirely free from perturbations by diurnal and other species of tide. The values change from day to day, and computed quantities are precisely defined at a certain fixed hour of the day. Essentially therefore,  $P$  and  $Q$  determine what we may regard as the average tide for the day. The observational values not only show the regular variations to be expected from the normal tidal constituents but they also show erratic day to day changes, obvious on account of their magnitude. It is clear, therefore, that the method is of very great value, even if the analysis for the tidal constituents is not affected, in providing observational data for meteorological perturbations of the semi-diurnal tides (and similarly of all other species of tide and of sea level). If, however, the analysis is completed so that the harmonic constituents are determined, then it is possible to synthesize these constituents (say for the semi-diurnal species only) to give synthesized values of  $P$  and  $Q$  which we can call  $P_s$  and  $Q_s$ . Alternatively the whole of the data can be expressed in the form  $R$ ,  $\epsilon$ ,  $R_s$ , and  $\epsilon_s$ .

It was the examination of the quantities

$$\lambda = R/R_s \quad \text{and} \quad \eta = \epsilon - \epsilon_s,$$

which led originally to the present investigation.

The synthesized semi-diurnal tide will include all the principal tidal constituents, and thus  $\lambda$  and  $\eta$  will reveal—

- (i) Harmonic constituents neglected in the synthesis,
- (ii) Harmonic constituents of unknown origin,
- (iii) Erratic variations, probably due to meteorological changes.

Further details of the method are given in Appendix I.

### 3 *Perturbations of the Semi-diurnal Tide*

The method already described was applied to the semi-diurnal tide at Liverpool for the year 1918. Values of  $R$  and  $\epsilon$  had already been tabulated

at the time of the analysis, whilst  $R$ , and  $\tau$ , were obtained from quantities generated on the predicting machine from the 1918 Harmonic Constants

Daily values of the quantities  $\lambda$  and  $\eta$  are given in Table I

A cursory examination of the results shows that in general the perturbations occur more or less casually, though on several occasions they remain noticeably consistent over a period of several days. The maximum perturbation is about 15 minutes in times and 1.5 feet in range. It seems reasonable to suppose that eventually the casual variations will be explained in terms of the meteorological conditions. Up to the present, however, correlation methods which have been applied have shown little success. This would further indicate, 'that in the ultimate solution of the meteorological tide, dynamical methods must largely replace the statistical methods of the past'

Table I also contains monthly averages of the quantities  $\eta$  and  $\lambda$  which we shall call  $\eta_m$  and  $\lambda_m$

Monthly time errors of the synthesized tide are indicated by  $\eta_m$ . It will be seen that in April the average time error amounted to 3 minutes, whilst in the remaining months of the year it was 2 minutes or under. There is no obvious periodicity in the values, but it is noticeable that in April and again in August, when the time errors were greatest, the range was nearest normal

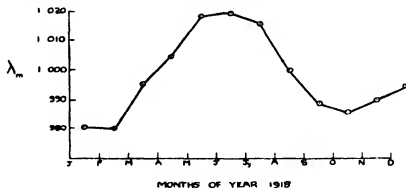


FIG. 1—Annual perturbation in the range of semi-diurnal tide at Liverpool, 1918

Monthly range errors are indicated by  $\lambda_m$ . These show a prominent perturbation which is illustrated in fig. 1. It is clear that the actual tide is less than the synthesized tide during the winter months and greater than the synthesized tide during the summer months, the discrepancy in range being of the order of 2%. With an average semi-range of 10 feet this indicates an average discrepancy of nearly 3 inches in semi-range of tide during the summer months.

Table I—Daily Perturbations  $\eta$  and  $\lambda$ [illegible]

### Monthly Perturbations $\eta_m$ and $\lambda_m$

$\lambda_{90}$	$\lambda_{80}$	$\lambda_{70}$	$\lambda_{60}$	$\lambda_{50}$	$\lambda_{40}$	$\lambda_{30}$	$\lambda_{20}$	$\lambda_{10}$	$\lambda_0$	$\lambda_{-10}$	$\lambda_{-20}$	$\lambda_{-30}$	$\lambda_{-40}$	$\lambda_{-50}$	$\lambda_{-60}$	$\lambda_{-70}$	$\lambda_{-80}$	$\lambda_{-90}$					
0.7	0.981	0.0	0.980	-0.7	0.997	1.5	1.046	0.0	0.019	0.7	1.020	0.7	1.016	1.1	0.999	-0.1	0.988	-0.7	0.960	-0.7	0.990	0.7	0.983



The existence of a perturbation of this magnitude, whether casual or continued, is of considerable practical importance quite apart from any theoretical interest it may possess, and it is with a more complete investigation of the annual portion of this perturbation that the present paper is concerned. If the phenomenon were due to a single harmonic semi diurnal constituent of amplitude 0.2 feet we should expect it to disturb high and low water times by 2 minutes, but the observed disturbance of times is not as great as this. We shall show later that the disturbance can be expressed by means of two harmonic constituents.

#### *4 Annual Perturbation of the Principal Constituent of Semi-diurnal Tide*

Tidal analysis in the past has been restricted to a search for constituents indicated by the tide generating potential and by the shallow water theory. In the Report of the British Association for 1923 Proudman has shown that additional frictional constituents are possible.

None of the above suggests an annual variation in the range of tide.

For the purpose of continuing the investigation, a direct method of analysis for such an annual perturbation was considered desirable, and in Appendix II Doodson's tables for the analysis of tidal observations have been extended to include two new constituents which have been called  $MA_2$  and  $Ma_2$ , and which perturb  $M_2$ , the principal lunar constituent, once in the year. These constituents, which are conjugates with respect to  $M_2$ , have been chosen so that  $MA_2$  loses and  $Ma_2$  gains on  $M_2$  very approximately 1 degree per day.

For comparison purposes a method is also given whereby the constants obtained for  $MA_2$  and  $Ma_2$  may be combined so as to give an expression for the annual perturbation in semi-range of  $M_2$  in the form

$$R \cos (\alpha + 1^\circ d),$$

where

$R$  = the amplitude of the perturbation (semi-range),

$\alpha$  = the phase lag of the perturbation on January 1,

$d$  = the number of days elapsed from January 1

#### *5 Analytical Results at Liverpool*

Five years' observations of the Liverpool tides, 1918, 1920, 1922, 1924 and 1930, have been analysed for the new constituents, and the results are given in Table II.

We may summarize our observations on this table as follows —

(i) There can be little doubt of the continued existence at Liverpool of an annual perturbation of  $M_2$ .

(ii) The perturbation shows a definite consistency of phase from year to year. In all five years the range of phase lies within  $55^\circ$ , whilst if we exclude the year 1924 the range is within  $30^\circ$  (A phase difference of  $30^\circ$  corresponds to a time difference of 1 month.)

Table II — Analytical Results at Liverpool

Year	$MA_1$		$Ma_1$		$R \cos(\alpha \pm 1^\circ d)$		$\% \frac{H(MA_1)}{H(M_2)}$	$\% \frac{H(Ma_1)}{H(M_2)}$	$\% \frac{R}{H(M_2)}$
	H	$\phi$	H	$\phi$	R	$\alpha$			
1918	0.123	217	0.102	20	0.198	196	1.22	1.01	1.98
1920	0.118	230	0.069	289	0.087	220	1.17	0.69	0.87
1922	0.127	228	0.033	305	0.115	195	1.26	0.33	1.05
1924	0.045	273	0.057	342	0.098	248	0.45	0.57	0.98
1930	0.193	324	0.173	93	0.166	222	1.90	1.71	1.66
Vector average	0.086	256	0.040	24	0.126	213	0.86	0.40	1.25

(iii) In 1918 the perturbation was unusually large, amounting to nearly 2% of  $M_2$ . In 1920 it was less than half this value. Over the five years examined the average perturbation was 1.25% of  $M_2$ .

A rather interesting fact is that the year to year magnitude would appear to follow the year to year magnitude of  $M_2$ .

(iv) If we confine our attention to the constituents,  $MA_1$  would appear to be the more important of the two, with an amplitude averaging double that of  $Ma_1$ . There is also an apparent tendency for the phase lag of  $MA_1$  to increase steadily with a period of roughly 40 years. This tendency was first noticed in the years 1918 to 1924 and is further confirmed by the year 1930. If further observations show this steady increase to be maintained, it will mean that the true period of the phenomenon is not exactly 1 year but slightly less.

It is convenient here to compare the results obtained from the monthly averages of  $\lambda$  with those from the direct analysis of hourly heights. In order to do this the twelve mean values of  $\lambda$  were analysed, and the results expressed in the form  $R \cos(\alpha \pm 1^\circ d)$ . The actual expression obtained from the analysis was

$$0.212 \cos(197 \pm 1^\circ d),$$

as compared with

$$0.198 \cos (196 + 1^\circ d),$$

from the 1918 analysis of hourly heights

These results show conclusively that the annual perturbation in range originally found in the semi-diurnal tide may be considered to be adequately represented by these two new constituents

### 6 *Perturbations in the High and Low Water Range of Tide*

In the preceding sections an examination was made first of the semi-diurnal tide as a whole, and then of the principal constituent of the semi-diurnal tide. We shall now extend this examination to the tide as observed at the times of high and low water.

The simplest method of proceeding would be to analyse the monthly averages of the differences between observed and predicted tides. A more satisfactory method is to make use of existing analyses for shallow water constituents.

[The method of analysis for shallow water constituents, though devised by Doodson several years ago has not yet been published. The quantities analysed are the daily differences between observed and predicted times and heights of high and low waters. A search for annual (Sa) variations in these quantities is included in the method. Simple differences between the Sa term for high water, and that for low water, give the perturbation in the range of tide. The semi-range of the perturbation may then be easily expressed in the form  $R \cos (\alpha + 1^\circ d)$ .]

One year's observations, 1924, have been analysed for Liverpool by the above method, and the expression obtained for the annual perturbation was

$$0.110 \cos (228 + 1^\circ d),$$

as compared with

$$0.098 \cos (248 + 1^\circ d),$$

from the analysis of 1924 hourly heights.

The main difference, that of  $20^\circ$  in phase, is not considerable, and could easily arise from the independent methods of analysis, one of hourly heights, and the other of high and low waters. It is possible, however, through variable shallow water effects, to have an annual perturbation in the high and low waters, independent of one in the semi-diurnal tide. This may also partially account for the difference.

### 7 *Annual Perturbation in British Waters*

(1) *From Hourly Heights*—Proceeding to the region of the British Isles generally, Table III gives the values of the constants  $MA_2$  and  $Ma_2$ , and the

deduced perturbation  $R \cos (\alpha + 1^\circ d)$  at all places in these regions for which analyses of hourly heights already existed at the Tidal Institute. Additional quantities are given in which amplitudes are expressed as a percentage of the amplitude of  $M_2$ . These quantities are also given in the tables for Liverpool, and are later used for comparison purposes. They will be referred to as "relative amplitudes."

Table III—British Waters, Annual Perturbation from Analysis of Hourly Heights.

Place	Year	$MA_1$		$M\alpha_1$		$R \cos (\alpha + 1^\circ d)$		$\% \frac{H(MA_1)}{H(M_2)}$	$\% \frac{H(M\alpha_1)}{H(M_2)}$	$\% \frac{R}{H(M_2)}$
		H	$\varphi$	H	$\varphi$	R	$\alpha$			
Stornoway	1920	0 020	57	0 055	153	0 038	321	0 44	1 20	0 83
Aberdeen	1930	0 030	261	0 016	221	0 038	133	0 70	0 35	0 89
Immingham	1926	0 139	13	0 160	159	0 076	223	1 39	2 19	1 04
Harwich	1931	0 073	276	0 045	113	0 082	196	1 72	1 06	1 92
Southend	1925	0 181	330	0 077	182	0 141	201	2 75	1 17	2 15
Southampton	1924	0 063	198	0 021	106	0 093	147	1 42	0 47	2 08
Portland	1924	0 055	151	0 011	231	0 061	245	2 67	0 53	2 70
Avonmouth	1924	0 108	161	0 084	200	0 180	257	0 77	0 60	1 79

The tables indicate at once that the perturbation exists generally in British waters, and at a number of places is of even greater importance than at Liverpool. An examination of the changes from place to place, in phase and amplitude of the constituents and perturbation, lead to the following interesting results.

Commencing at Stornoway the phase lag of  $MA_1$  steadily increases as one proceeds southwards down the North Sea, reaches a maximum at Immingham, and then diminishes slightly at Harwich and Southend. In the lower approaches it increases as one proceeds into the Irish Sea and also up the English Channel. This increase in phase lag as one advances from the Atlantic into the neighbouring seas is precisely what one would expect, were the constituent generated in the open sea, and propagated in a manner similar to that of an ordinary astronomical tide.

The changes of phase of  $M\alpha_1$  are not nearly so definite as those of its conjugate constituent. Over the whole of the North Sea and English Channel the phase is, within a range of 100 degrees, more or less constant. At Liverpool, in the Irish Sea, it differs by roughly 180 degrees from that in the above two regions.

Concerning the variations in the relative amplitudes of the constituents, of the two,  $MA_1$  is clearly the more important, though there are occasions, as at Stornoway and Immingham, where the reverse is true. Over the southern

North Sea and English Channel, the relative amplitude of  $MA_2$  is roughly 2% of  $M_2$ , and it is striking that this is more than double its value in the Irish Sea and on the Scottish coasts

The constituent  $Ma_2$  is practically constant over the Irish Sea and English Channel, where its relative amplitude is approximately 0.5% of  $M_2$ . In the North Sea its amplitude is somewhat greater, and at Immingham rises to a prominent maximum of 2% of  $M_2$ .

If we confine our attention to the resultant annual perturbation the indications are, with the possible exception of one station, Aberdeen, that the perturbation occurs steadily later in the year as one proceeds inward from the Atlantic

At the approaches, both North and South, the maximum perturbation occurs between the middle and end of February. At Southend and Harwich it occurs in June, whilst at Liverpool it occurs about half a month earlier.

If we take an average of  $\alpha$  over the whole region we get a value of  $214^\circ$ . If we take an average of the corresponding phase for the change in sea level, we get a value of  $76^\circ$ . The difference between the two is roughly  $140^\circ$ . This might suggest a tendency for the perturbation in range to be opposite in phase to the perturbation in sea level. This would mean a maximum perturbation in range to be associated with a minimum sea level, and *vice versa*.

As to the changes in the relative amplitude of the perturbation, in the Irish Sea and English Channel the perturbation would appear to decrease in amplitude as one proceeds inwards from the Atlantic. In the North Sea an opposite result is found. Between Stornoway and Immingham the perturbation is practically constant, whilst a very rapid increase occurs at Harwich, and this is further maintained at Southend.

The importance of the perturbation relative to  $M_2$  is at Immingham and Aberdeen 1% of  $M_2$  and at all other places 2% of  $M_2$ .

(II) *From High and Low Waters*—Table IV gives the expressions for the annual perturbation as derived from existing shallow water analyses. For the majority of places the results are in fair agreement with those from hourly heights. It is noticeable that where agreement is least, particularly in phase, shallow water constituents are abnormally large.

It will be noticed that up to the moment no mention has been made of London Bridge. This was unnecessary because of the close agreement of the results with those of Southend.

The stations are, however, worth comparing one with another, because of light they may throw on variations in the perturbation in a large tidal river.

Comparisons between yearly amplitudes and phases at the two places show that abnormal values at the one are associated with corresponding abnormalities at the other, whilst the five year average indicates the perturbation at London Bridge to be slightly greater, and to occur a little later than at Southend. It

Table IV—British Waters, Annual Perturbation from Harmonic Shallow Water Analysis.

Place	Year	$R \cos (\alpha + 1^{\circ} d)$		$\frac{R}{H} (M_s)$
		R	$\alpha$	
Harwich	1931	0.085	181	2.00
Southend*	1925	0.085	230	1.30
	1926	0.054	224	0.83
	1927	0.101	181	1.54
	1928	0.095	225	1.45
	1931	0.151	188	2.30
	Vector average	0.061	205	1.39
London Bridge*	1925	0.077	201	0.96
	1926	0.074	218	0.93
	1927	0.148	178	1.86
	1928	0.078	215	0.98
	1931	0.141	192	1.77
	Vector average	0.097	199	1.22
Southampton	1924	0.070	252	1.58
Portland	1924	0.058	301	2.80
Avonmouth*	1924	0.150	284	1.07
	1925	0.108	257	0.78
	1926	0.172	249	1.23
	Vector average	0.137	263	0.99
Liverpool	1924	0.110	228	1.09

\* The vector average refers only to Southend, London Bridge and Avonmouth.

is difficult from the results to imagine the origin of the perturbation to be in the river, the indications are much more in favour of it being propagated inwards from the mouth.

### 8 Perturbation in General

Table V contains the results of analyses, at a small number of places, widely distributed over the surface of the globe.

*St John, N B*, is situated in the Bay of Fundy in a region where resonance effects may be expected. The results both in amplitude and phase are very similar to those at certain places in the British Isles, in particular to Harwich and Southend. Two years of analyses exist and it is clear, as in previous cases, that considerable variation in amplitude, and to a less degree in phase, can occur from year to year. It is interesting that the results further confirm what was previously found to exist at Liverpool, viz., that yearly changes in

Table V—Perturbation in General

Place	Year	$MA_1$		$Ma_2$		$R \cos (\alpha + 1^\circ d)$		$\frac{1}{100} \frac{H(MA_1)}{H(M_1)}$	$\frac{1}{100} \frac{H(Ma_2)}{H(M_2)}$	$\% \frac{R}{H(M_2)}$
		H	$\phi$	H	$\phi$	R	$\alpha$			
St. John, N B	1930	0 174	283	0 024	158	0 162	246	1 75	0 24	1 03
"	1931	0 231	265	0 094	152	0 296	168	2 30	0 94	2 95
Ponta Delgada	1924	0 054	248	0 049	324	0 013	222	3 48	1 15	0 84
Johore Bahru	1929	0 071	178	0 020	113	0 081	152	2 55	0 72	2 90
Port Hedland	1927	0 047	317	0 047	58	0 050	229	0 82	0 82	0 87
Newchwang	1924	0 352	196	0 203	190	0 322	150	0 20	5 30	8 40

the amplitude of the perturbation are accompanied by yearly changes in the amplitude of  $M_2$ . Thus in

1930  $H$  of  $M_2 = 9 957$  whilst  $R = 0 162$

1931  $H$  of  $M_2 = 10 083$  whilst  $R = 0 296$

*Ponta Delgada*, in the Azores is suitably situated to be typical of tides in the middle of the Atlantic. It is very striking that the magnitude of the perturbation found here should be practically the same as at Stornoway and Aberdeen on the open Scottish coasts.

*Port Hedland*, which is placed on the North-west coast of Australia, is in an open position comparable with Ponta Delgada, but on the opposite side of the globe. The magnitude of the perturbation at the two places is much the same.

*Johore Bahru*, in the Singapore Strait, has a prominent perturbation larger than at any place in the British Isles. The conditions at this place are peculiar, for it is situated in a comparatively deep channel, but the flow of water through the channel is restricted by an artificial causeway. The winds along the direction of the channel have an annual variation.

*Newchwang*, on a tidal river in the Yellow Sea, is icebound for a portion of the year. As a result, considerable freshet effects are experienced at times, and this makes the results of particular importance. The existence of a large perturbation in the range of tide at Newchwang has been known for a number of years, and it has always been attributed to the very abnormal conditions. What is of importance is that practically the whole of the perturbation exists in the semi-diurnal tide, and does not arise, except to a very minor extent, from variations in the shallow water constituents.

### 9 *Annual Perturbation and Local Meteorological Conditions*

It was stated early in the paper, that little success had resulted from an attempt to correlate the day to day changes in the perturbation with the local meteorological conditions. It is now proposed to replace day to day values by mean values for the month. Also an assumption (similar to that adopted by Doodson when considering mean sea level variations\*) will be made that the average perturbation in range of tide can be expressed as a linear function, of the departure from the mean of the local pressure, and of the local east and north pressure gradients. Thus if

$B$  = the departure from the mean of the average local pressure for the month,

$E$  = the departure from the mean of the average local easterly pressure gradient for the month,

$N$  = the departure from the mean of the average local northerly pressure gradient for the month

then the average range perturbation for the month ( $\lambda_m$ ) =  $\alpha B + \beta E + \gamma N$ , where  $\alpha$ ,  $\beta$ ,  $\gamma$ , are constants depending upon the correlation and also on the units used.

Twelve mean monthly values of  $\lambda_m$  at Liverpool were obtained from the curve for the average perturbation, for the years 1918, 1920, 1922, 1924, 1930.

Monthly averages for  $B$ ,  $E$ ,  $N$ , at Liverpool were obtained from "The Weather of the British Coasts M.O. 230". The figures are based upon the 7-hour observations at the Telegraphic Reporting Stations of the Meteorological Office in the ten years 1906 to 1915.

With special reference to Liverpool

$B$  = the monthly average of the departure from the mean of the pressure at Liverpool,

\* 'Mon. Not. R. Astr. Soc., Geophys. Suppl.', vol. 1, p. 124 (1924)



E = the monthly average of the departure from the mean of the pressure between Spurn Head and Blacksd Point,

N = the monthly average of the departure from the mean of the pressure between Thurso and Plymouth

These quantities are given in Table VI The twelve equations obtained from the above values were solved by the method of least squares for  $\alpha$ ,  $\beta$ ,  $\gamma$  The results gave

$$\lambda_m = 0.7B - 8.0E + 2.8N$$

Fig 2 shows the annual perturbation, together with the perturbation as deduced from the meteorological conditions by substitution in the above formula

Table VI—Value of B, E, N, at Liverpool from 10 year Average

	Jan	Feb	Mar	Apr	May	June	July	Aug	Sept	Oct	Nov	Dec
B	1.6	-3.2	-2.5	1.8	1.8	2.3	2.0	0.8	5.3	-1.2	-2.2	-7.3
E	0.8	0.4	-1.0	0.1	0.3	-1.1	-1.5	-0.7	-0.2	2.4	-0.8	+1.4
N	-4.3	-4.3	-0.9	0.9	3.3	2.4	1.4	0.5	1.9	3.7	-1.5	-2.6

The annual variation is undoubtedly reproduced by the meteorological conditions, though there are marked differences in many of the monthly values. A tendency for the results from the meteorological conditions to lag roughly one month behind the actual perturbation is hard to explain. It is possibly due to the selection of local meteorological conditions instead of those far out at sea.

The contributions from B, E, N, to the perturbation in fig 2 as deduced from the meteorological conditions, were in the ratio 21.86.70. From this we would infer the effects from easterly pressure gradients to be slightly greater than those from northerly pressure gradients, and roughly four times greater than the effect from local barometric pressure.

### 10 Previous Experience of an Annual Perturbation

The earliest reference to a seasonal change in semi-diurnal tide is to be found in Darwin's paper on the "Harmonic Analysis of the Antarctic Tidal Observations of the 'Discovery'." The observations discussed in his paper were taken during the years 1902 and 1903 at a position on the south-eastern

extremity of Ross Island. For analytical purposes the observations were grouped into periods of one month. The feature of these monthly analyses which was of particular interest was that the  $M_2$  tide showed a progressive annual change both in amplitude and phase during each of the two seasons analysed. The change was such that the average range was least and the average tide most retarded during the months of June, July and August. The magnitude of the change was even greater than at any of the places discussed in the present paper, having a relative amplitude of 38%. It is particularly interesting that no other constituent showed this progressive annual change. Darwin also found that an inequality of 13 months would

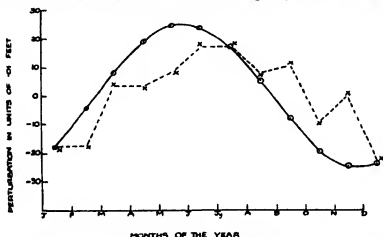


FIG. 2.—Perturbation as deduced from meteorological conditions. Perturbation, — actual, - - - as deduced from meteorological conditions.

satisfy the conditions better than one of 12 months and from this he suggested a possible connection with the 14 months period for the variation in latitude.

In recent years several references to an annual perturbation in the range of tide have appeared in special publications of the US Coast and Geodetic Survey. The references have been based upon monthly averages of the observed high and low waters at individual places. In Special Publication No. 162, "On the tides and currents in Chesapeake Bay and Tributaries," we find the following remarks—"At Philadelphia on the Delaware River, the average seasonal variation in range is about 1 foot, whereas at Boston near the open coast it is only 0.2 feet. It appears, therefore, that such a variation is largely a river effect, and it is probably due to the varying quantities of drainage water in the river." The existence of the perturbation at Newchwang was discovered in 1925 following a series of monthly analyses made at the

Tidal Institute The perturbation was so prominent and so regular that it has been included in the Newchwang predictions since 1927

Analyses for shallow water constituents were commenced in 1924 and the unequal annual changes in the high and low waters were obvious from that time The small number of these analyses available prevented an examination of the perturbation at an earlier date

### 11 Discussion

Perhaps the most fundamental of all the causes likely to produce a perturbation in the range of tide, is a periodic change in the boundary conditions This cause will not provide a universal explanation of the phenomenon, but it should be particularly applicable to the seas of the polar regions Darwin found that in the Antarctic the least range and greatest retardation of tide occurred during the months June, July and August, when the Antarctic seas are frozen over to their greatest extent When we consider that it is during these months that the greatest restrictions to the free flow of the currents will be present, the occurrence of a reduced range of tide during this period is exactly what we should expect

In a tidal river the perturbation may arise in several ways, provided there is an annual change in the average depth or in the non-tidal current in the river A change in the average depth from the ordinary law for the propagation of a wave in a canal will affect the speed and consequently the phase at a particular place to a first order, whilst it will affect the range to a second order The phase and range of tide will also change with any variation of the non-tidal current in the river This may be seen as follows —

The equations of motion for the tides in a canal of rectangular section will be found in Lamb's "Hydrodynamics" If we neglect second order terms other than those arising from the interaction of tidal and non-tidal current and consider the currents as made up of two parts, one a tidal portion  $u$  and the other a non-tidal portion  $U$ , the modified equations of motion become

$$\begin{aligned}\frac{\partial u}{\partial t} + g \frac{\partial \zeta}{\partial x} &= -U \frac{\partial u}{\partial x} \\ \frac{\partial \zeta}{\partial t} + h \frac{\partial u}{\partial x} &= -U \frac{\partial \zeta}{\partial x},\end{aligned}$$

where  $x$  is the distance from the origin along the length of the canal,  $g$  is the acceleration due to gravity,  $t$  is the time,  $\zeta$  is the elevation of the surface from the mean level.

Solving these differential equations in the usual manner and assuming a tidal oscillation of the form  $Re^{i\sigma t}$ , where  $\sigma$  is the speed, the expression for  $\zeta$  becomes

$$\zeta = Ae^{i\left[\frac{\sigma x}{c-U} + t\right]} + Be^{i\left[\frac{\sigma x}{c+U} - t\right]},$$

where  $A$  and  $B$  are constants and  $c^2 = gh$

The real part of this may be written in the form  $\zeta = R \cos(\sigma t - \epsilon)$ , where

$$R^2 = A^2 + B^2 + 2AB \cos \frac{\sigma x}{c^2 - U^2}$$

$$\tan \epsilon = - \frac{A \sin \frac{\sigma x}{c-U} - B \sin \frac{\sigma x}{c+U}}{A \cos \frac{\sigma x}{c-U} + B \cos \frac{\sigma x}{c+U}}$$

Clearly, then, provided  $A$  and  $B$  are neither equal to zero, any change in  $U$  will produce a corresponding change in  $R$  and  $\epsilon$

The range of tide in a river may also be affected by changes in certain of the shallow water constituents which themselves may vary from changes in the average depth and of the constant current

The observational data already discussed in this paper have shown that though in particular cases, as in the Delaware River, the changes can largely be explained as a river effect, this is not universally so, and the explanation is not applicable to the River Thames where the perturbations at Southend and London Bridge are found to be practically the same

The formula, deduced above for a tidal oscillation and a variable non-tidal current, is also applicable to an open ocean, provided we can prove the existence of a variable non-tidal current. The existence of such a current in the Atlantic having an annual variation, is now a well-established fact in oceanographical research, and is confirmed by the sea level changes in the marginal seas and by observations of sea temperature, salinity, density, distribution and character of plankton, both in the Atlantic Ocean, and in the marginal seas, and also in the connecting channels. Clearly, however, from the relations between  $c$  and  $U$  in the open ocean the changes in  $U$  must be much greater than can be reasonably expected if they are to produce the changes that are observed. Again, the present paper has shown that in the open ocean the magnitude of the perturbation is of the same order as the change in the mean sea level. This would practically rule out any explanation of the perturbation in terms of sea level changes in the open ocean. We

must admit then, that no satisfactory explanation of the perturbation in the open ocean has yet been found

### *Conclusion.*

We have seen established an annual perturbation in the range of tide, which at a number of places is of sufficient importance for inclusion in future tidal predictions. Under what conditions is the additional labour, which such an inclusion would involve, justified?

Considering the yearly variations in the perturbation, the results have shown, that though the consistency in phase can generally be relied upon, the value of the amplitude may vary considerably from year to year. This would lead us to proceed with caution where the amplitude is concerned, and it would appear advisable for this reason alone, to have two, and possibly three, years' analyses available, previous to considering the inclusion of this perturbation in predictions. A reasonable limit for the amplitude of the perturbation, would be one greater than 1% of that of  $M_2$ , though for places of particular consistency the limit might easily be fixed at a lower value.

As is necessary in a paper of this type, attention has been drawn to indications which later observations may, or may not, prove to be true. It is intended that these be taken, purely for what they are worth, on the information available, and that they be kept in mind as future data come to hand. More definite and satisfying conclusions should be possible by a revision, and expansion of the paper, in, say, ten years' time.

### *Acknowledgments.*

The investigation has been carried out, at the original suggestion, and under the personal direction, of Doodson, to whom the writer wishes gratefully to acknowledge his indebtedness for suggestions and advice. Aid in checking the computations and analyses was willingly provided by the Tidal Institute Staff. Certain information on general oceanography was made available by the late Professor Johnstone.

### APPENDICES

In the following appendices constant reference will be made to Doodson's paper on the "Analysis of Tidal Observations," and it will be assumed that the reader is familiar with both the notation and theoretical basis of this paper.

## APPENDIX I

Let

$\zeta$  = the tidal elevation above a known datum

$\sigma$  = the increment in phase of a constituent, in degrees per mean solar hour

$\rho$  = the increment in phase in degrees per mean solar day with integral multiples of  $360^\circ$  omitted

$t$  = the number of hours measured from the origin of time

$T$  = the number of complete days measured from the origin in time

$R$  = amplitude of the constituent

$R \cos(\sigma t - \varepsilon)$  = a typical tidal constituent with phase  $-\varepsilon$  at the origin of time

$H$  = the hour of the day measured from midnight

Taking a tidal constituent in the form

$$R \cos(\sigma t - \varepsilon),$$

the height of tide due to this constituent at hour  $H$  on the day  $T$  may be written in the form

$$R \cos(\sigma H + \rho T - \varepsilon)$$

If we take any combination of hourly heights such as  $\zeta_H + \zeta_{H+1} + \zeta_{H+2}$  then it can be shown that the results may be expressed in the form

$$JR \cos(\sigma H + \rho T - \varepsilon + \eta),$$

where  $J$  and  $\eta$  are constants depending upon the combination used

Doodson has shown that it is possible to obtain daily expressions of the above form which are functions of one species only. For the semi-diurnal tide two such expressions are obtained

$$X = JR \cos(\rho T - \varepsilon + \eta')$$

$$Y = JR \cos(\rho T - \varepsilon + \eta'')$$

The quantities  $\eta'$  and  $\eta''$  are chosen so that the initial hour of  $X$  is midnight preceding the day, whilst that of  $Y$  is 3 hours later

The computation of the initial phase of the amplitude require two further expressions

$$A = X + Y$$

$$B = X - Y,$$

of the form

$$A = aR \cos(\delta - \rho T) \quad \text{and} \quad B = bR \sin(\delta - \rho T),$$

so that

$$a = 2J \cos \frac{1}{2}(\eta' - \eta'') \quad \text{and} \quad b = -2J \sin \frac{1}{2}(\eta' - \eta''),$$

$$\delta = \epsilon - \Delta \quad \text{and} \quad \Delta = \frac{1}{2}(\eta' + \eta'')$$

It will thus be seen that for any particular day the phase of both A and B—

(1) as obtained from observations through X and Y,

(2) as obtained by generation from the final constants, at intervals of 24 hours from the time origin (taken as zero hour of the central day),

is the phase of the tidal constituent (1) from observation, (2) from synthesis,  $\Delta/\sigma$  hours after the preceding midnight

The value of  $\Delta/\sigma$  for the average semi-diurnal tide is 15.5 hours. Thus if on any day we write the semi-diurnal tide in the form

$$R \cos(30^\circ t - \epsilon') = P \cos 30^\circ t + Q \sin 30^\circ t,$$

where  $t$  = the time in solar hours measured from 15.5 hours after the preceding midnight,  $-\epsilon'$  = the phase of the semi-diurnal tide at this time

$$P = R \cos \epsilon' \quad \text{and} \quad Q = R \sin \epsilon',$$

then

$$\frac{b}{a} \tan \epsilon' = \frac{\Sigma B \text{ (for the semi-diurnal constituents)}}{\Sigma A \text{ (for the semi-diurnal constituents)}},$$

and

$$\Sigma(a^2 + b^2) R^2 = \Sigma A^2 + \Sigma B^2$$

To a first approximation the magnitude of both  $a$  and  $b$  for the semi-diurnal tide may be taken as 34

The values of  $R$  and  $\epsilon'$  from observation are easily found through  $\Sigma X$  and  $\Sigma Y$  the quantities tabulated daily in the analysis. Values of  $R$ , and  $\epsilon$ , the synthesized quantities, are obtained from  $\Sigma A$ , and  $\Sigma B$ , which are generated on the predicting machine. The quantities  $R$  and  $\delta$  necessary in the synthesis are obtained from the completed analysis, whilst values of  $a$  and  $b$  are given in Doodson's paper.

The L  g   predicting machine which was used for the purpose provides for ten semi-diurnal constituents as follows —

$$2SM_2, 2N_2, \mu_2, N_2, \nu_2, M_2, L_2, T_2, S_2, K_2.$$

Since we are only interested in daily increments of the constituents it is possible

to generate additional minor constituents by making use of the following relationships

- MNS<sub>2</sub> has the same  $\rho$  as MN<sub>4</sub>
- R<sub>2</sub> has the same  $\rho$  as K<sub>1</sub> (very approx.)
- MSN<sub>2</sub> has the same  $\rho$  as Mm
- OP<sub>2</sub> has the same  $\rho$  as Mf but of opposite sign
- $\lambda_2$  has the same  $\rho$  as Mm but of opposite sign

The constituents MSN<sub>2</sub> and  $\lambda_2$  were generated as compound constituents.

Much care was taken in the reading of  $\Sigma A_2$  and  $\Sigma B_2$ . The scale used was 4 cm to 1000 units, readings being to the nearest 0.05 cm.

## APPENDIX II.

### *Analysis Tables for an Annual Perturbation of M<sub>2</sub>*

Constituents which will produce an annual perturbation of M<sub>2</sub> may be considered as having either of the following arguments

$$[(\text{Argument of } M_2) - \lambda]$$

or

$$[(\text{Argument of } M_2) + \lambda],$$

where  $\lambda$  is the mean longitude of the sun

In the present section Doodson's analysis tables have been extended to include the above two constituents which have been called MA<sub>2</sub> and Ma<sub>2</sub>, respectively

The eight new functions which it is necessary to calculate for the determination of the new constituents are in the notation of Doodson's paper, for

$$R \cos \delta = A_{221}, B_{220}, B_{201}, A_{200},$$

and for

$$R \sin \delta = B_{221}, A_{220}, A_{201}, B_{200}$$

Table VII contains values of  $\sigma$ ,  $\rho$ ,  $\Delta$ ,  $a$ ,  $b$ ,  $D$ ,  $M$ , for MA<sub>2</sub> and Ma<sub>2</sub>. The quantities  $\sigma$  and  $\rho$  are the hourly and daily speeds of the constituents, whilst  $\Delta$  is a constant used in determining the final constants,  $a$ ,  $b$ ,  $D$ ,  $M$ , are necessary for the calculation of the number of contributions from MA<sub>2</sub> and Ma<sub>2</sub> to the various functions

Table VIII sets out in tabular form the correction process for  $[R \cos \delta]$  and  $[R \sin \delta]$  whereby contributions from all constituents other than MA<sub>2</sub> and Ma<sub>2</sub> are eliminated. Except for a correction through the terms  $A_{121}$ ,



and  $B_{126}$  for an annual perturbation which arises from  $O_1$ , and also the use of a uniform divisor of 10,000, the tables are similar in form to those of Doodson. Table IX contains the factors for the calculation of  $R \cos \delta$  and  $R \sin \delta$  from  $[R \cos \delta]$  and  $[R \sin \delta]$

Table VII.—Data relative to  $MA_2$  and  $Ma_2$

	$MA_2$	$Ma_2$
$\sigma$	28 943036	29 025173
$\rho$	-25 367146	-23 396851
$\Delta$	88 5	89 9
$a_2$	35 276	35 146
$b_2$	33 377	-33 446
$D_a$	-1 462	2 024
$D_b$	-2 862	3 488
$D_c$	29 964	30 476
$D_d$	91 574	30 152
$D_e$	2 342	-0 832
$D_f$	-1 220	0 092
$M_a$	0 457	0 372
$M_b$	11 272	11 216
$M_c$	-14 020	15 124
$M_d$	-0 464	0 274

Table VIII  
Correction Tables for the Calculation of  $[R \cos \delta]$

Principal function	Correction terms—Multiples of										Divisor	Symbol
	$A_{201}$	$A_{202}$	$A_{211}$	$B_{210}$	$A_{220}$	$A_{221}$	$B_{220}$	$A_{230}$	$A_{231}$	$A_{232}$		
$A_{201}$	0 004	0 002	-0 023	0 039	-0 009	0 026	-0 073	0 003	-0 010	0 083	10,000	A
$B_{201}$	-0 005	0 001	0 081	-0 018	0 005	-0 140	0 007	—	0 012	-0 040	10,000	B
$A_{211}$	0 084	-0 023	-0 029	0 067	-0 008	0 041	-0 040	0 001	0 008	0 029	10,000	C
$A_{220}$	0 109	0 012	-0 160	0 016	-0 006	0 086	-0 039	—	0 012	0 094	10,000	D

Correction Tables for the Calculation of  $[R \sin \delta]$

Principal function	Correction terms—Multiples of										Divisor	Symbol
	$B_{101}$	$A_{101}$	$B_{211}$	$A_{210}$	$B_{220}$	$B_{221}$	$A_{230}$	$B_{230}$	$B_{231}$	$B_{232}$		
$B_{201}$	0 004	—	-0 022	-0 038	-0 009	0 027	0 082	0 002	-0 009	0 040	10,000	A'
$A_{201}$	0 005	0 002	-0 081	-0 018	-0 005	0 164	0 007	-0 001	-0 010	0 130	10,000	B'
$A_{211}$	-0 084	-0 019	0 031	0 067	0 009	-0 049	-0 040	-0 002	-0 006	-0 029	10,000	C'
$B_{220}$	0 109	-0 004	-0 161	-0 015	-0 006	0 086	0 039	0 001	0 006	0 045	10,000	D'

*Constituents expressed as a Perturbation in Range of  $M_2$ .*

Let  $H_1$ ,  $H_2$ ,  $H_3$  be the amplitudes and  $g_1$ ,  $g_2$ ,  $g_3$  the phase lags of  $M_2$ ,  $MA_2$ ,  $Ma_2$ , respectively

Then the phases of the constituents relative to  $M_2$  are, of  $MA_2$

$$(g_1 - g_2 - h),$$

and of  $Ma_2$

$$(g_1 - g_3 + h)$$

Table IX —Calculation of  $R \cos \delta$  and  $R \sin \delta$  from  $[R \cos \delta]$  and  $[R \sin \delta]$

Constituent	R cos $\delta$				R sin $\delta$			
	Multiples of—				Multiples of—			
	A.	B	C	D	A'	B'	C'	D'
$MA_2$	0 41	-0 32			-0 43	-0 30		
$MA_2$			0 48	0 34			0 45	-0 35
$Ma_2$	0 41	0 34			-0 42	0 32		
$Ma_2$			0 42	-0 32			0 40	0 35

The perturbation in the semi-range of  $M_2$  is given by

$$[H_2 \cos \{h - (g_1 - g_2)\} + H_3 \cos \{h + (g_1 - g_3)\}],$$

which may be expressed in the form

$$R \cos (h + \beta),$$

where

$$\tan \beta = \frac{H_2 \sin (g_1 - g_2) - H_3 \sin (g_1 - g_3)}{H_2 \cos (g_1 - g_2) + H_3 \cos (g_1 - g_3)},$$

and

$$R^2 = [H_2^2 + H_3^2 + 2H_2H_3 \{\cos (2g_1 - g_2 - g_3)\}]$$

If  $\alpha$  is the phase of the constituent on January 1 when the value of  $h$  is  $280^\circ$ , and if  $d$  are the number of days elapsed from January 1, then since the daily increment in  $h$  is with sufficient accuracy equal to  $1^\circ$  per day, an alternative form for the annual perturbation is

$$R \cos \{(\alpha + 280 + \beta) + 1^\circ d\}$$

This is the form used in the paper

*Summary*

The paper establishes the existence of an annual perturbation in the range of tide. The perturbation became evident during an intensive examination, by a method which should be of particular interest, of the residual semi-diurnal tide at Laverpool. Direct analysis of hourly heights for an annual perturbation in the principal constituent of semi-diurnal tide showed the perturbation to be consistent from year to year, and to exist generally in British waters and throughout the world at large. Analysis of high and low waters further confirmed the results. A study of the distribution of the perturbation provided a number of interesting results as also did a study of its relation with local meteorological conditions. The paper concludes with a short discussion on possible causes of an annual perturbation in the range of tide. Appendices describe the method by which the residual tide was obtained, and also contain tables for the direct analysis of hourly heights for an annual perturbation in the range of tide.

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*The Production of Showers by Cosmic Radiation*

By C W GILBERT, B A

(Communicated by Lord Rutherford, O M, F R S—Received November 17, 1933)

*Introduction*

The production of secondary particles by the passage of cosmic radiation through matter is a well-known feature of the cosmic ray phenomenon. Cloud chamber photographs of Blackett and Occhialini,\* Anderson,† and Kuntze‡ show that the secondaries often occur in groups, which have been named "showers". The production of these showers was investigated by observing the simultaneous discharges of three Geiger-Müller counters so arranged that they only recorded showers of at least three particles. The main experiments were carried out at the Forschungs-station Jungfraujoch, in Switzerland, at a height of 3500 metres. Other experiments were made at Eigergletscher.

\* 'Proc Roy Soc,' A, vol. 139, p 699 (1933)

† 'Phys. Rev,' vol. 43, p 368 (1933), vol. 44, p 406 (1933)

‡ 'Z Physik,' vol. 80, p 559 (1933)

(2300 m.) and Zürich (500 m.) to investigate the variation of frequency of the showers with altitude

### *Experimental Method*

The experimental arrangement was similar to that used by Rossi\* and by Fünfer† Three Geiger-Müller counters were used, arranged as shown in fig. 1 The lead under investigation was placed above in such a position

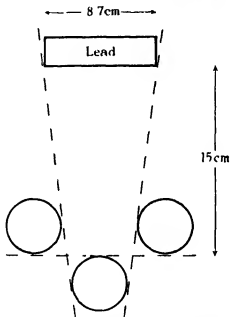


Fig. 1 — Experimental arrangement of counters and lead.

that no particle coming from the lead could go through more than one counter. The triple coincidences due to the presence of the lead must thus be due to showers of at least three particles. The coincidences were recorded by a method described by Rossi‡. The impulse from a fourth valve was fed to the grid of a thyratron by resistance capacity coupling, fig. 2. This impulse was a *negative* kick, but the accompanying positive back-kick was sufficient to trip the thyratron. This arrangement rendered the circuit very stable and free from external pickup.

\* 'Z. Physik,' vol. 82, p. 151 (1933).

† *Ibid.*, vol. 83, p. 92 (1933).

‡ 'Nature,' vol. 125, p. 636 (1930).

The counters used were 20 cm long  $\times$  4.2 cm diameter. These gave about 120 counts per minute at sea level and between 200 and 300 per minute at 3500 m, at this rate the mechanical telephone counter used could not respond to all the counts. The number of casual triple coincidences was only 2 or 3 per hour. From the number of casual double coincidences the resolving power was found to be  $10^{-3}$  seconds. The rate of casual triple coincidences was very much smaller than the rate of counting of triple coincidences and so no correction was made for them.

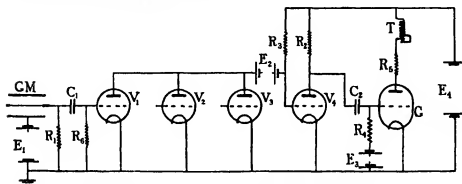


FIG. 2—GM, Geiger Müller counter, T, telephone counter and break,  $R_1$ ,  $4 \times 10^5 \Omega$ ,  $R_2$ ,  $20,000 \Omega$ ,  $R_3, R_4, R_5$ ,  $10^7 \Omega$ ,  $R_6$ ,  $800 \Omega$ ,  $E_1$ , about 1500 v,  $E_2$ , about 10 v,  $E_3$ , about 10 v,  $E_4$ , about 100 v

Difficulty was encountered at the Forschungs-station Jungfraujoch, on account of the damp atmosphere, and the roof of the building was so thick that satisfactory measurements were only possible on the roof. So the counters and recording set were placed in a tin box which stood on the roof, which was then protected by a wooden box from snow and sun. Leads were taken from the tin box through a length of large bore rubber tube into the building for the L T supply, counter HT, telephone counter, etc. The air in the tin box was dried with  $P_2O_5$  and the box heated by two 40-watt lamps. The voltage for the counters was supplied by a 1500-volt dry battery and it was found necessary to place this near a heater to keep it sufficiently dry at night. By these means it was possible to work the counters on the roof, day and night under all weather conditions\*.

The coincidences were recorded on a telephone counter and also on an automatic recorder supplied by the Cambridge Instrument Company. This recorded each coincidence on a slowly moving paper disc, and gave a useful

\* The mean atmospheric temperature in July was about  $1^\circ \text{C}$ . It frequently snowed and was often very misty.

check on the performance of the apparatus, as well as the necessary data for investigating the distribution of counts in time

### Experimental Results

Triple coincidences were observed at Jungfraujoch with the arrangement shown in fig 1, with varying thicknesses of lead over the counters. As the thickness of lead was increased the number of coincidences was found to rise to a maximum at 2.2 cm. and then to fall off rapidly (Table I and fig 3). The curve

Table I—Dependence of showers on the thickness of lead

Thickness of lead in cm	0.0	0.77	1.35	1.90	2.19	2.48	2.87	3.69	5.77	7.67	9.46
Number of periods of counting	7	7	6	8	4	3	4	6	5	4	3
Total time in hours and minutes	11 01	8 28	7 49	13 39	7 31	7 14	10 27	5 33	6 46	6 15	6 44
Total count	559	545	562	1069	614	560	791	383	393	336	386
Mean count per hour	50.9	64.5	72.0	78.4	81.9	77.9	75.6	69.1	58.0	53.8	57.3

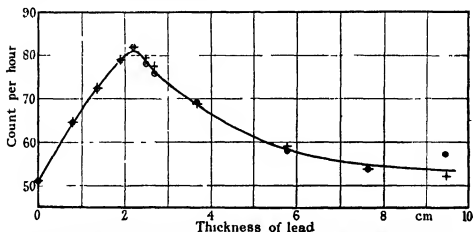


FIG 3—Showers observed with different thicknesses of lead above the counter at 3500 m. C, observed points, +, calculated.

is very similar to curves obtained by Rossi and Fünfer at sea level. The most notable difference is the different position of the maximum, which for Rossi's and Fünfer's curves was at 1.6 cm.

With no lead triple coincidences are obtained at a greater rate than the rate of casual coincidences, which are due to showers originating in the atmosphere

This background count will not be greatly affected by the presence of the lead. After subtracting this background the fall away from the maximum is exponential with a coefficient of  $\mu = 0.38 \text{ cm}^{-1}$ . Rossi's and Fünfer's curves gave rather lower values. The curve will be discussed in more detail later.

Fünfer observed that the lead side screens round his counters gave an increased count which was dependent on the count from the lead over the counters. He found that the maximum effect was reached with a thickness of the side screens of 0.6 cm. He obtained a similar effect with lead under the counters\*. To investigate this 2.0 cm of lead were placed over the counters and varying thicknesses placed underneath, an increase in the counting rate was obtained which reached a maximum at 0.6 cm, fig. 4. Then 2.0 cm of

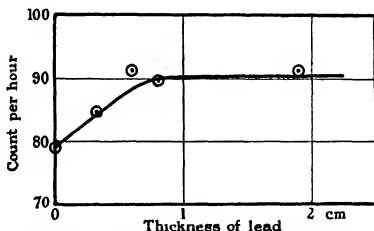


Fig. 4.—Showers observed with different thicknesses of lead under the counter

lead were placed under the counters and the lead above varied. The curve obtained, fig. 5, was similar to the one without lead underneath. The difference between the curves of figs. 3 and 5 also shows a maximum, fig. 6. Because of the statistical errors the position of this maximum cannot be fixed at all accurately, but it almost certainly lies not far from 2 cm. The maximum increase is about 40% of the count due to the upper lead. Fünfer had attributed this effect to reflection by the lower lead of the shower particles from the upper lead. However, such an explanation would require that about 20% of the shower particles would be reflected back through nearly 180°. Cloud chamber photographs show that there is no such large backward scattering. Further, Blackett and Occhialini obtained conclusive evidence of high

\* 'Z Physik,' p. 83, vol. 92 (1933)

energy particles thrown backwards from showers. The explanation thus appears to be that the increase is due to particles thrown in a backward direction from showers occurring in the lower lead.

Measurements were made at Jungfraujoch, Eigerletscher\* and Zürich† to determine how the increase with altitude of the frequency of the showers

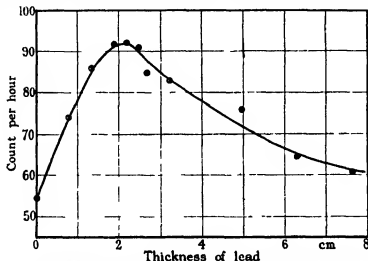


FIG. 5—Showers observed with 2.0 cm of lead under the counter and different thicknesses above.

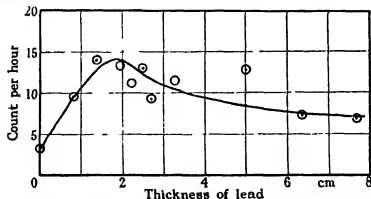


FIG. 6—Showers due to lead under the counter with different thicknesses of lead over the counter. Difference between curves of fig. 3 and fig. 5

\* I wish to thank the management of the Hotel Eigerletscher for allowing me to carry out these measurements on the hotel roof and for their kind assistance.

† I also wish to thank Professor Dr. P. Scherrer of the Physikalisches Institut der Technischen Hochschule, Zürich, for his kind permission to make the Zürich measurements on the roof of the Institute. I am also very grateful for the assistance given me by the staff.



compared with the increase of the general cosmic radiation. At these places measurements were made with the arrangement shown in fig 1 with 2.2 cm of lead over the counters, the frequency of showers being taken as the difference in counting rate with and without the lead. Measurements were also made with the three counters one above the other in a vertical plane. This gave a measure of the general cosmic ray intensity. Since the maximum of the showers at Zurich is below 2.2 cm, the frequency of showers for this station must be corrected. This is done by estimating the number, using the curve of fig 3, that would have been recorded if the shower particles had had sufficient energy to reach the counters. The results are shown in Table II.

Table II—Comparison of frequency of showers and cosmic radiation intensity

	Jungfrauoch	Figergletscher	Zürich
Atmospheric pressure, in cm. of Hg	50.7	57.0	72.3
Showers from 2.2 cm. of lead, count per hour	31.0	24.3	11.6
Corrected	—	—	14.4
Counters in vertical plane, count per minute	5.45	4.40	2.50
Ionization intensity, ions per c.c. per sec.	5.68	4.21	2.74
Showers	1.00	0.78	0.46
Cosmic radiation intensity—			
Counters	1.00	0.80	0.45
Ionization	1.00	0.73	0.48

The first line gives the showers observed from 2.2 cm. of lead, the second line the rate of counting with the counters in a vertical plane. The third line shows the cosmic ray intensity measured by Wollan with an ionization chamber\*. The next three lines are the same measurements reduced so that the value at Jungfrauoch is 1.00. The number of showers thus increases with altitude at about the same rate as the general cosmic ray intensity, as measured by both the total ionization and the number of particles producing coincidences in these three counters placed in a vertical plane.

#### *The Transition Curve for Lead*

The curve showing the dependence of the number of showers recorded on the thickness of lead above is very similar to those obtained by Rossi and by Fünfer. The chief difference is the change of the position of the peak. The

\* Compton, 'Phys. Rev.', vol 43, p 394 (1933)

position of the maximum observed at Jungfraujoch was 2.2 cm. Both Rossi and Fünfer located the maximum at 1.6 cm. Preliminary measurements made at Cambridge before going to Jungfraujoch had shown the peak to occur at about 1.6 cm, and later measurements made at Cambridge confirm this, fig. 7.

It appears certain that the curve must be interpreted\* to be a transition effect on passing from air to lead. The shower particles to which the counters respond must be charged since they produce ionization in the counters, and so they have a range in lead†. When the thickness of lead is greater than the range of the shower particles the showers from the upper part of the lead can

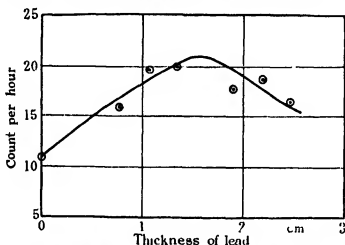


Fig. 7—Showers observed with different thicknesses of lead above the counters at sea level.

no longer be recorded. Hence the curve will have a maximum at the thickness of lead equal to the range of the shower particles in lead. The curve must then fall off with the absorption coefficient of the radiation producing the showers. Since this absorption coefficient is very much larger than the absorption coefficient of the general radiation,  $0.38 \text{ cm}^{-1}$  in lead as compared with  $0.006 \text{ cm}^{-1}$ , the immediate cause of the showers cannot be the general cosmic

\* Rossi, 'Nature,' vol. 132, p. 173 (1933), see also Johnson's "Analyses of Schindler's transition curves," 'Phys. Rev.,' vol. 41, p. 545 (1932).

† A charged particle loses energy continually and so has a definite maximum range. But the particles of an uncharged radiation lose all or at least a considerable portion of their energy in a single event, so the radiation is absorbed exponentially. There is, therefore, a finite probability that some of the radiation will travel any given distance and have no maximum range.

radiation Since the absorption coefficient of the radiation producing the showers is so large and also the absorption exponential, the production of showers is probably the main cause of absorption We shall call this radiation the shower-producing radiation This radiation could not penetrate the whole of the atmosphere and so must be produced by some other radiation, which we shall call the primary radiation After traversing sufficient depth of homogeneous matter an equilibrium will be reached between the amount of primary and shower producing radiation present The curves obtained are thus transition curves due to the readjustment of this equilibrium in passing from one medium to another

Suppose that the shower producing particles are produced by a primary radiation with absorption coefficient  $\mu$  and so the intensity of the primary at any depth  $x$  in the atmosphere is  $I = I_0 e^{-\mu x}$ , treating the problem as one-dimensional, where  $x$  is measured downwards

Let  $I_s$  be the intensity of the shower-producing radiation at depth  $x$  and let  $\mu_s$  be the absorption coefficient, and  $\alpha$  the coefficient of production of shower-producing radiation by the primary  $\alpha$  may be considerably less than  $\mu$  since the production of shower-producing radiation may not be the only means of absorption of the primary

Then

$$\frac{dI_s}{dx} = -\mu_s I_s + \alpha I$$

The solution is

$$\begin{aligned} I_s &= B e^{-\mu_s x} + \frac{\alpha}{\mu_s - \mu} I_0 e^{-\mu x} \\ &= B e^{-\mu_s x} + A I_0 \end{aligned}$$

on writing

$$A = \frac{\alpha}{\mu_s - \mu}$$

If  $\mu < \mu_s$ , then after traversing a large thickness of matter the second term is the only significant one, and so the equilibrium intensity of the shower-producing radiation is given by

$$I_s = A I_0$$

$A$  is a constant for any homogeneous medium but may vary from medium to medium.

Consider now the effect of putting varying thicknesses of a material over the counters, see fig 8

Let

$S$  be the number of showers observed

$a$  the mean range of the shower particles in the material

$Z$  the thickness of material whose upper surface is at  $x = 0$

Let the index (1) denote coefficients in the air and index (2) coefficients in the material

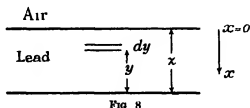


FIG. 8

We have

$$I_p^{(2)} = B e^{-\mu_p^{(2)} z} + A^{(2)} I$$

But at  $x = 0$

$$I_p^{(2)} = I_p^{(1)} - A^{(1)} I,$$

and so

$$B = (A^{(1)} - A^{(2)}) I$$

We thus have, where  $\beta = z$  if  $z < a$  and  $a$  if  $z \geq a$ ,

$$\begin{aligned} S &= C \int_0^\beta \mu_p^{(2)} I_p^{(2)}(z-y) dy, \\ &= C \int_0^\beta \mu_p I_0 \{ (A^{(1)} - A^{(2)}) e^{-(\mu_p + \mu)(z-y)} + A^{(2)} e^{-\mu(z-y)} \} dy, \\ &= \begin{cases} C(A^{(1)} - A^{(2)}) I_0 \frac{\mu_p}{\mu_p + \mu} (1 - e^{-(\mu_p + \mu)\beta}) + C A^{(2)} I_0 \frac{\mu_p}{\mu} (1 - e^{-\mu\beta}) & \text{for } z \leq a, \\ C(A^{(1)} - A^{(2)}) I_0 \frac{\mu_p}{\mu_p + \mu} (e^{(\mu_p + \mu)a} - 1) e^{-(\mu_p + \mu)z} + C A^{(2)} I_0 \frac{\mu_p}{\mu} (e^{\mu a} - 1) e^{-\mu z} & \text{for } z \geq a, \end{cases} \end{aligned}$$

where  $C$  is a constant depending on the chance of observing a shower. To this number of showers must be added the background. Thus, in the experimental arrangement of fig. 1, will be approximately constant. Denote this background by  $D$ . If  $\mu < \mu_p$ , then the second term has little effect on the shape of the curve until a considerable thickness has been reached. Thus for small thicknesses the shape of the curve is given by

$$S = \begin{cases} C'(1 - e^{-\mu_p z}) + D & \text{for } z \leq a \\ C'(e^{\mu_p z} - 1) e^{-\mu z} + D & \text{for } z \geq a \end{cases}$$

This formula agrees very well with the experimental results with the values

$$a = 2.2 \text{ cm}$$

$$\mu = 0.38 \text{ cm}^{-1}$$

$$C' = 53.9 \text{ counts per hour}$$

$$D = 51.0 \text{ counts per hour}$$

The points calculated using these values are shown by crosses in fig. 3.

For large thicknesses of material the tail of the curve falls away less rapidly and finally has an absorption coefficient  $\mu^*$ . The sharpness of the peak of the experimental curve suggests that the energy of the shower particles is fairly homogeneous†. The curves of Rossi and Finfer have not such sharp peaks. This may be due to the fact that they used much wider sheets of lead and so many showers would be recorded that had come through the lead at an angle.

The shape of this transition curve is similar to the transition curves (Übergangskurven) obtained by Schindler using an ionization chamber‡. The fall of his curves immediately past the maximum gives an absorption coefficient which agrees well with the values obtained with counters. The position of the maximum is also in rough agreement. It thus appears certain that the transition effects observed by Steinke,§ Schindler (*loc cit*) and others with ionization chambers are essentially the same as those observed with counters, and therefore are mainly an effect due to the showers.

### The Shower Particles

The position of the peak of the curve varies from 1.6 cm. at sea level to 2.2 cm. at 3500 m. The corresponding energies of the shower particles as calculated from Heisenberg's|| formula are 70 and 95 million electron volts, if the particles are positive and negative electrons. The measurements of Anderson¶ confirm Heisenberg's data at these high energies. That the particles are positive and negative electrons seems certain from the expansion chamber photographs of Blackett and Occhialini (*loc cit*) and Anderson\*\*.

\* Rossi, 'Nature,' *loc cit*

† Bhabha, 'Z. Physik,' vol. 86, p. 120 (1933)

‡ Schindler, 'Z. Physik,' vol. 72, p. 625 (1931)

§ 'Z. Physik,' vol. 48, p. 647 (1928)

|| 'Ann. Physik,' vol. 13, p. 430 (1932)

¶ 'Phys. Rev.,' vol. 43, p. 381 A (1933), vol. 44, p. 406 (1933)

\*\* 'Phys. Rev.,' vol. 44, p. 406 (1933).

Fünfer's (*loc cit*) results give the ranges of the shower particles of showers produced in lead, iron and aluminium as 1.6 cm, 2.5 cm and 7.0 cm in lead, iron and aluminium respectively. All these values correspond to an energy of approximately 70 million electron volts. Thus it appears that the energy of the shower particles does not vary greatly with the atomic number of the material.

#### Shower Producing Radiation

The absorption coefficient of the shower producing radiation as observed at Jungfraujoch is  $0.38 \text{ cm}^{-1}$  in lead. Since the absorption coefficient is exponential the absorption probably takes place through single, or at the most a few, events. Thus the greater part of the absorption is probably due to the production of showers. The relation

$$\mu = N\sigma,$$

where  $N$  is the number of nuclei per unit volume and  $\sigma$  the effective cross-section for production of a shower, gives the value of cross-section for this absorption coefficient as  $11.4 \times 10^{-24} \text{ sq cm}$  (radius  $1.90 \times 10^{-12} \text{ cm}$ .) This is a very large cross-section, being larger than the nucleus itself. From the data of various workers the cross-section for production of a shower can be estimated for lead, iron and aluminium. The results are shown in Table III. Thus

Table III — Effective cross section for production of a shower

	Lead	Iron	Aluminium.
Absorption coefficient of shower producing radiation in $\text{cm}^{-1}$ of the material	0.30* 0.29† 0.38‡ 0.34§	0.064* 0.11† — —	0.017* — — —
Radius of cross-section $\times 10^{12} \text{ cm}$ .	17.5	5.8	3.0
Atomic number, $Z$	82	26	13
Area cross-section $\frac{\sigma}{Z^2} \times 10^{27}$	1.43	1.54	1.66

\* Fünfer, *loc cit*.

† Rossi, 'Z. Physik,' vol. 82, p. 151 (1933)

‡ The author

§ Schindler, 'Z. Physik,' vol. 72, p. 628 (1931)

the area of cross-section varies nearly as the *square* of the atomic number. This is in striking contrast to the usual mass absorption rule holding for cosmic radiation in general. Recent calculations of Oppenheimer and Plesset||

|| 'Phys. Rev,' vol. 44, p. 53 (1933)

show that for  $\gamma$ -rays of high energy the cross-section for production of a pair of positive and negative electrons is given by

$$\sigma \sim \frac{Z^2}{137} \frac{e^4}{m^2 c^4}$$

Furry and Carlson\* show that for high energy electrons the cross section is of the same order. For lead the cross-section derived by Oppenheimer and Plesset is  $3.9 \times 10^{-24}$  sq cm as compared with the observed value of  $11.4 \times 10^{-24}$  sq cm for the absorption coefficient of the shower-producing radiation †

The mechanism of production of the showers by the shower-producing radiation is probably that pairs of electrons are produced in the field of the nucleus. Oppenheimer and Plesset have only dealt with the production of a single pair, but it is possible that in the showers many pairs are produced simultaneously. The showers thus appear to be creation of particles and not, as at first looked probable, the disintegration of the nucleus.

The experiment with lead under the counters shows that there are showers with particles thrown backward with a range in lead of approximately 0.6 cm that is an energy of about 30 million electron volts. Further, these showers depend in some way on the showers produced in the lead above, for as the thickness of the lead above is increased the number of showers from the lower lead increases, reaches a maximum lying between 1 and 3 cm. and then decreases.

Thus the showers from the upper lead must contain some kind of radiation which is capable of producing showers in the lower lead. This is in agreement with the remark of Blackett and Occhialini (*loc cit*, p. 717) that "when one shower occurs there is a surprisingly large chance that another will occur a short way below it. This leads one to the view that a shower-producing radiation can be produced in the showers themselves, but as is shown later, this cannot be the main cause of the shower-producing radiation."

About the nature of the shower-producing radiation little can be said. Its chief means of absorption is probably the production of showers and since the mean free path for this is so small no information is given by its absorption as to whether it is charged or not. It may possibly consist of gamma radiation.

\* 'Phys. Rev.', vol. 44, p. 237 (1933).

† Heitler and Santer, 'Nature,' vol. 132, p. 892 (1933), obtain a cross section for production of positive and negative electrons in lead of  $33 \times 10^{-24}$  sq cm for  $\gamma$  rays of energy 50 million electron volts.

*Primary Radiation*

Since the shower-producing radiation has such a small mean free path, there must be some mechanism of reproducing the radiation. We have seen that it can be reproduced in the showers themselves, but this cannot be the only method for this alone cannot give the transition effects observed. These require that the shower producing radiation be in equilibrium with some other radiation and that the equilibrium value changes on passing from one material to another. It thus appears necessary that the shower-producing radiation is produced by a primary radiation which traverses the whole atmosphere.

For large thicknesses of material the number of showers falls off with the absorption coefficient of the primary radiation. Further, the number of showers increases with altitude at the same rate as the general radiation. So the absorption of the primaries is probably the same as that of the general cosmic radiation. Now the absorption of the general radiation in large thicknesses of different material obeys approximately the mass absorption rule. Hence the primary radiation probably obeys the same rule. From the absorption measurements of cosmic radiation in large thicknesses of material we can get an upper limit for the cross-section of *production* of the shower-producing radiation by the primary radiation (Table IV).

Table IV — Effective cross-section of absorption of primary radiation

	Lead	Iron	Water
Absorption coefficient in $\text{cm}^{-1}$ of the material	0.0064*	0.0070*	0.00021†
Area of cross section $\times 10^{18} \text{ cm}^2$	2.5	1.7	0.44

\* Schindler, 'Z. Physik,' vol. 72, p. 628 (1931)

† Regener, 'Phys. Z.,' vol. 34, p. 306 (1933)

The majority of the particles present in the cosmic ray stream in the atmosphere may have originated in showers. The primaries would then probably be charged so as to give the observed latitude variation.

*Energy Relations*

The fact that the energy of the showers increases upwards in the atmosphere suggests that the energy of the showers comes from the primary radiation and is not released from the nucleus by some trigger action.

The energy carried by the shower particles is probably of the order of  $1.5 \times 10^6$  electron volts at sea level, assuming 20 particles in a shower. Since



the energy required for an electron to penetrate the atmosphere is  $5 \times 10^9$  electron volts, the energy of the primary radiation is probably of the order of  $10^{10}$  electron volts. Particles of this order are required by the theory of Lemaître and Vallarta\* to explain the observed latitude effect. On going from 3500 m. to sea level an electron which had  $10^{10}$  electron volts energy when it entered the atmosphere will suffer a 26% decrease in energy. This compares favourably with the 25% decrease of the energy of the shower particles between these levels.

I am indebted to the Department of Scientific and Industrial Research for a grant to enable me to work in the Cavendish Laboratory, and to the Royal Society for a grant to cover the travelling expenses of myself and the apparatus to Switzerland for the measurements at high altitudes.

I wish to thank Professor W. R. Hess and the administration of the Hochalpine Forschungs-station, Jungfrauoch for permission to make use of that station, and for their kind assistance.

I wish also to thank Professor P. M. S. Blackett for his invaluable help both in the conduction of the experiments and in the discussion of the results.

#### Summary

Experiments made with triple coincidence counters show that the frequency of showers produced in lead by the passage of cosmic radiation is proportional to the general cosmic radiation. The transition curves for air to lead were obtained at 3500 m. and it was found that there the energy of the shower particles was greater than at sea level. To explain the curves obtained, three types of radiation are needed, a primary radiation, a shower-producing radiation and the shower particles. The nature and properties of these radiations are summarized in Table V.

Table V

Radiation	Kind	Evidence	Absorption.
Primary	Probably charged	Latitude effect	Mass absorption $\sigma \approx 7$
Shower producing	Unknown		Exponential absorption $\sigma \sim \frac{Z^2}{137} \frac{e^4}{m^2 c^4}$
Shower particles	Positive and negative electrons	Wilson cloud chamber photographs	Mass absorption range $\sim 1/\rho$ .

\* 'Phys. Rev.', vol. 42, p. 914 (1932)

# *The Nuclear Spin of Tin*

By S. TOLANSKY, Ph.D. (1931 Exhibition Senior Student), Imperial College of Science, London

(Communicated by A. Fowler, F.R.S. - Received November 22, 1933)

## *Introduction*

In a previous note to 'Nature,'† the author has briefly reported the main conclusions arrived at from an investigation of the fine structures of the lines of the spark spectrum of tin. It was stated there that the nuclear spin of the two main odd isotopes of tin was  $\frac{1}{2}$ , the magnetic moments being negative and also identical in value for both. Almost immediately after this note appeared, an independent report was published by Schüler and Westmeyer,‡ who reached identical conclusions about the spins. The present paper gives a detailed account of the author's observations on the Sn II spectrum. In addition to this, the value of the nuclear spin has been confirmed by measurements in the Sn I spectrum.

Aston has reported 11 isotopes in tin, the relative abundances of which are shown in Table I.

Table I

Isotope	112	114	115	116	117	118	119	120	121	122	124
% abundance	1.1	0.7	0.4*	14.2	9.8*	21.5	11.0*	27.0	2.9*	5.0	6.2

The odd isotopes, indicated by asterisks, comprise about 24% of the total. In accordance with what has usually been observed in atoms containing a mixture of odd and even isotopes, it is to be expected that the even isotopes will have zero nuclear spin, or at least very small nuclear magnetic moments. Each even isotope will contribute a single component to the pattern of a given line, and if there is no isotope displacement effect these will all superpose, producing a single strong component.

On the other hand, the odd isotopes will possess nuclear mechanical and magnetic moments, and the coupling of the nuclear magnetic moment with the external electron magnetic moment of the atom will cause the terms to split, such that the even isotope in each term is at the centre of gravity of the split

† 'Nature,' vol. 132, p. 318 (1933).

‡ 'Naturwiss.,' vol. 21, p. 690 (1933).

term. As a result, the expected line patterns will consist of tiny multiplet groups, the summed intensity of which will be 24% of the total, and at the centre of gravity of each group will be a single strong line comprising 76% of the total intensity, and representing the contribution of the even isotopes. If this central strong component is split up or even appreciably broadened, this is an indication that even isotope displacement is taking place.

The present investigation is largely concerned with the Sn II lines emitted by a hollow cathode discharge in tin. The term scheme of this spectrum has been given by Green and Loring\* and further details have been reported by Lang†.

The transitions which have been studied here are shown in the term scheme in fig. 1. Other transitions lie in a region unsuited to the silvered Fabry-Pérot interferometer which was used as the resolving instrument. In the normal state, the tin ion has a  $5s^2 5p$  electron configuration, so that a group of doublet terms is built up by the  $5p$  electron going into various orbits. From the point of view of fine structure analysis, the closed  $5s^2$  group can be neglected and the doublet terms treated as if they arise from a one electron system. Such a procedure is entirely justified both on theoretical grounds and by previous experimental evidence. However Lang (*loc cit*) has shown that in addition to the doublet group discussed above, there exists a doublet and quartet system based upon  $5s 5p^2$ . The observed terms due to this configuration are shown to the right of fig. 1 and it will be shown later that there is evidence that fine structure perturbations take place between the  $^3D$  terms of the two configurations.

### Experimental

The Sn II spectrum was excited both in water cooled and hot hollow cathode discharges. The tubes employed were slight modifications of the type used so successfully by Schüler for the production of intense lines free from broadening effects. Helium was continuously circulated through the tubes at about a millimetre pressure and currents of 200–300 milliamperes employed. The resulting tin spectrum was extremely bright. The lines were examined for fine structure by means of a variable gap, silvered, Fabry-Pérot interferometer of large aperture, crossed with a two prism spectrograph. Using Ilford hypersensitive plates the pattern of the strongest line could be photographed in a few minutes, although it includes a relatively weak component whose intensity is only 5% of that of the complete pattern.

\* 'Phys. Rev.', vol 30, p. 574 (1927)

† 'Phys. Rev.', vol 35, p. 445 (1930)

The observed structures are shown in Table II. As usual, the separations are given in  $\text{cm}^{-1} \times 10^{-3}$ , and the visual estimates of intensity indicated below each component in brackets.

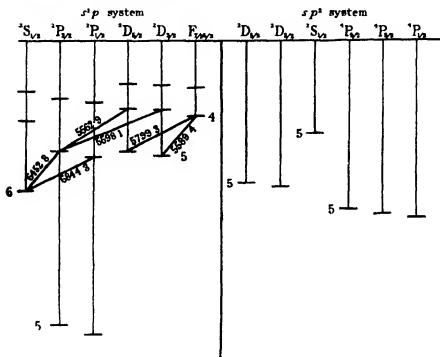


FIG. 1

Table II

Wave length	Allocation	Structure	Total width.
			$\text{cm}^{-1} \times 10^{-3}$
6444 3	6s $^3P_{1/2}$ —6p $^3P_{1/2}$	+ 100 65 0 181 (1) (3) (15) (1)	281
6462 8	6s $^3P_{1/2}$ —6p $^3P_{3/2}$	+ 83 0 154 — (3) (15) (1)	207
5799 3	5d $^3D_{3/2}$ —4f $^3F$	+ 152 0 239 — (3) (15) (2)	391
5598 1	6p $^3P_{3/2}$ —6d $^3D_{3/2}$	Single—sharp	—
5589 4	5d $^3P_{3/2}$ —4f $^3F$	Single—diffuse	~ 75
5562 9	6p $^3P_{3/2}$ —6d $^3D_{3/2}$	Single—sharp	—
5545 9*	?	+ 115 54 0 81 (2) (3) (6) (1)	250
4511 5*	?	+ 69 0 186 — (1) (2) (2)	255

\* It is not quite certain that these two lines belong to the Sn II spectrum

*Analysis of Structures.*

The analysis of each line will now be separately dealt with, and it will be shown that all the structures are completely accounted for on a nuclear spin of  $\frac{1}{2}$  for the two main odd isotopes, and a zero spin (or minute nuclear magnetic moment) for the even isotopes. The lines will be considered in the order best suited for analysis.

$\lambda 6452.8$  ( $6s\ ^2S_{1/2}-6p\ ^2P_{3/2}$ )—The structure in this line consists of a strong central component contributing about 75% of the total intensity of the pattern, flanked on either side by two weak components whose intensity ratio is 3 : 1. It is at once apparent that the central strong component arises from the even isotopes and as the two odd isotopes 117 and 119 are present to the extent of 9.8% and 11.0% respectively, the two weak satellites are almost entirely produced by these, any contribution from 121 being difficult to observe since its components will only be of intensity 1% and therefore practically negligible. The 3 : 1 intensity ratio of the components shows that both isotopes contribute to each component. The ratio of the intervals between the satellites and the central strong component is exactly 1 : 3, so that the strong line is at the optical centre of gravity of the pattern. The lower term is a  $6s\ ^2S_{1/2}$  term, and as it involves a penetrating  $s$  electron it will show fine structure. On the other hand, it is already well known that the  $6p\ ^2P_{3/2}$  term should show a relatively very minute structure, for all  $p\ ^2P_{3/2}$  terms yet investigated have conformed to this rule. It follows that the structure in the line is the structure of the lower term.

The number of fine structure levels in a term is  $2J + 1$  when the spin,  $I$ , is greater than  $J$ , or  $2I + 1$  when  $J > I$ . From this it is seen that a term with  $J = \frac{1}{2}$  always splits up into two fine structure levels, no matter what the value of  $I$ . An intensity ratio of 3 : 1 is only produced by a spin of  $\frac{1}{2}$ , the next value, namely,  $3/2$  giving an intensity ratio of 5 : 3 which could not be confused with the previous value. However, it will be shown later that the  $^2D_{5/2}$  term is also double, and as the  $J$  value is  $5/2$  here, this proves conclusively that the spin is  $\frac{1}{2}$ . In addition, it will be further shown that a spin of  $\frac{1}{2}$  is unambiguously deduced from fine structures in the Sn I spectrum.

The analysis for the line under consideration is shown in fig. 2. The upper term is single. The nuclear spin  $\frac{1}{2}$  combines with the lower  $J = \frac{1}{2}$  to give the fine structure quantum numbers  $F = 0$  and  $1$ . Both the odd isotopes must be treated as one, since otherwise four satellites instead of two would appear. The dotted lines show the even isotope transition, and in order to get this the

lower even isotope term has been placed at the centre of gravity of the lower term, *i.e.*, one-fourth of the distance from the level with  $F = 1$  (the quantum weight of a fine structure level  $F$  is  $2F + 1$ ). As the stronger satellite is to the violet, the fine structure levels are inverted, the higher  $F$  term lying deeper. This means that the nuclear moment is negative, so that  $I = -\frac{1}{2}$ . It will be observed that the calculated position of the even isotope line is 155 units

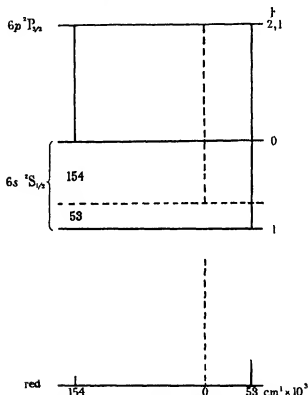


FIG. 2.—Fine structure of  $\lambda$  6452 8 ( $6s \ ^2S_{1/2} - 6p \ ^2P_{1/2}$ )

from the weaker component and experimentally it is found to be 154 units away. This is considered to be exact agreement.

$\lambda$  6844 3 ( $6s \ ^2S_{1/2} - 6p \ ^2P_{1/2}$ )—The structure of the lower term has already been determined from the analysis of the previous line, so that by simple analysis that of the upper term can be directly obtained. It was found experimentally that the intense even isotope line blurred the near by component of intensity 2, making accurate measurement of the position of this component very difficult. However, this does not invalidate the analysis, for it will be

seen from fig 3 that the structure of the upper term is determined by the separation of the remaining outer components. The even isotope level is again placed at the optical centre of gravity in both terms. The calculated structure is shown at A and the observed pattern at B. The agreement is

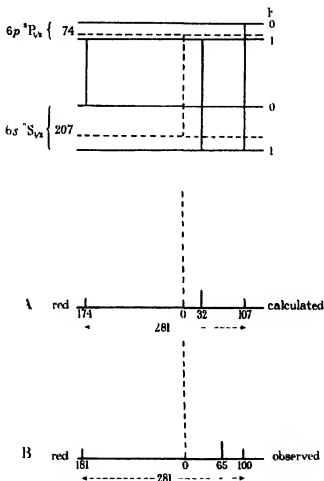


FIG. 3.—Fine structure of  $\lambda 6844$   $\frac{1}{2}$  ( $6s \ ^3S_{1/2} - 6p \ ^3P_{1/2}$ )

fairly satisfactory when account is taken both of the smallness of the separations and the difficulties caused by the over-exposed even isotope central component.

The upper term has a structure 74 units in width, with an uncertainty of perhaps 10 units at the utmost. As in the lower term the levels are inverted. The transition  $\Delta F = 0 \rightarrow 0$  is, of course, absent.

$\lambda 5598\ 1\ (6p\ ^2P_{3/2}-6d\ ^2D_{5/2})$ —This line shows no trace of structure, thus confirming the previous conclusion that the  $6p\ ^2P_{3/2}$  term is single. It proves that the upper term,  $6d\ ^2D_{5/2}$  also shows no structure.

$\lambda 5799\ 3\ (5d\ ^2D_{5/2}-4f\ ^2F)$ —There is a strong central component due to the even isotopes, but instead of being quite sharp, as in the previous lines, it is distinctly diffuse. In addition, there are two weak satellites arising from the odd isotopes, the structure being relatively quite wide, for the components are 391 units apart. The odd isotope structure is due entirely to the lower term, for it will be shown later that the  $4f\ ^2F$  term is single. This is also shown by —

- (a) The fact that the even isotope is almost at the centre of gravity of the doublet.
- (b) The fact that the stronger component is nearer to the violet shows that the structure is in the lower term, since the spin is negative.

The analysis of the line is shown in fig. 4. Of particular importance is the fact that the lower term has only two fine structure components, for this at once proves that the spin is  $-\frac{1}{2}$ .

The diffuse nature of the central component indicates a very small even isotope displacement.

$\lambda 5589\ 4\ (5d\ ^2D_{3/2}-4f\ ^2F)$ —This line is also single but is diffuse. The absence of weak components shows that the structure in both upper and lower odd isotope terms is extremely minute. This therefore confirms the previous conclusion that  $4f\ ^2F$  shows no structure. It is difficult to tell whether the small isotope displacement is in the lower or the upper term. An approximate estimate of the width of the line is 75 units, which would indicate a displacement between the even isotopes of the order of 20 units. This tentative value can only indicate the order of the effect.

$\lambda 5562\ 9\ (6p\ ^2P_{3/2}-6d\ ^2D_{5/2})$ —This line is very sharp and single, and as the lower term has already been proved to be single by two other lines, it follows that the upper  $6d\ ^2D_{5/2}$  term is also single.

Two other lines, namely,  $\lambda\lambda 5545\ 9, 4511\ 5$ , show peculiar structures, but it is not known whether they belong to the spectrum of Sn II. These are quite strong, however, and have not been identified with any probable impurity lines.  $\lambda 4511\ 5$  is particularly intense and is reported by Hemsalech as a weak tin line occurring in the spark. Although none of the remaining observers quoted in Kayser's 'Handbuch der Spektroskopie,' vol. 6, have observed this line, it appears with great intensity in the discharge tube employed. That



does not, however, show whether or not it is an impurity line. The structures in these two lines are quite different from those in all the other lines examined, for neither possesses a strong central even isotope component. If these lines do belong to tin, then they both show large even isotope displacement effects. In particular the intensities of the components of  $\lambda$  5545.9 agree fairly well with the abundance ratios of the even isotopes, when it is remembered that the odd isotope components will be distributed amongst them. If this is correct,

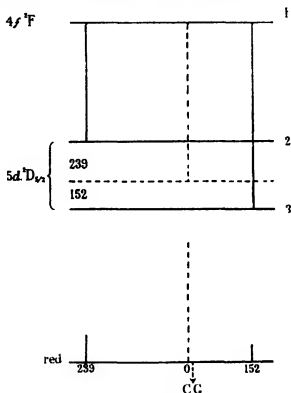


FIG. 4—Fine structure of  $\lambda$  5799.3 ( $5d^3D_{5/2} - 4f^3F$ )

it would imply an even isotope displacement of the order of 60 units. The peculiar structure in the other line could also arise from the superposition of a large isotope displacement effect on a small spin structure. It is, however, entirely unsafe to speculate about either of these lines since it is not known with certainty if they are tin lines or not.

### Discussion

It is seen that the fine structures are consistently explained by attributing a nuclear spin of  $-\frac{1}{2}$  to the two main odd isotopes. Both of these isotopes

have the same magnetic as well as the same mechanical moments, so that their respective patterns superpose exactly. There is no evidence of any measurable displacement effect between the two odd isotopes. That the spin must be  $\frac{1}{2}$  is proved by the structure of the  $5d \ ^3D_{5/2}$  term, for if  $I$  had any other value than this more than two components would have resulted. Furthermore the even isotope lines only fall at the centres of gravity of the patterns if the spin is taken to be  $\frac{1}{2}$ . As the higher valued  $F$  level lies deeper in all the terms, the spin is negative for both the isotopes, as for the neighbouring atom cadmium.

It is possible to deduce the interval factors in a number of terms, and these are shown in Table III. According to the interval rule, the interval between two fine structure levels  $F+1$  and  $F$ , is given by  $A(F+1)$  where  $A$  is a constant for a term and is called the interval factor. The values in the table refer only to the odd isotopes.

Table III

Term	Width of structure	Interval factor
$6s \ ^2S_{1/2}$	207	207
$6p \ ^2P_{1/2}$	74	74
$6p \ ^2P_{3/2}$	small	small
$5d \ ^2D_{5/2}$	391	130
$5d \ ^2D_{3/2}$	small	small
$6d \ ^2D_{5/2}$	small	small
$6d \ ^2D_{3/2}$	small	small
$4f \ ^3F$	small	small

In accordance with theoretical expectations, the  $6s \ ^2S_{1/2}$  term has a relatively wide structure, since it possesses a penetrating  $s$  electron. The structure in the  $6p \ ^2P_{3/2}$  term is too small to measure and this fact is in complete agreement with what has been found in other spectra. The interval factor for the  $6p \ ^2P_{1/2}$  term is 74. As in other one-electron spectra this is quite appreciable and appears to be due to the fact that in heavy atoms a  $p_{1/2}$  electron is relatively penetrating\*. Since the  $d$  and  $f$  electrons are non-penetrating, no observable fine structures are to be expected in terms involving them. It appears therefore that the interval factor 130 for  $5d \ ^2D_{5/2}$  is anomalously great. Associated with this is the question of the observed small even isotope displacement. This occurs in two lines  $5d \ ^2D_{5/2}-4f \ ^3F$  and  $5d \ ^2D_{3/2}-4f \ ^3F$ , but appears to be slightly more marked in the latter line, although this is difficult to decide.

\* Brent, 'Phys. Rev.', vol. 38, p. 462 (1931)

The multiplet separation of the  ${}^3F$  term is not resolved\*. It follows that the even isotope displacement occurs either in the two  ${}^3D$  terms or in the  ${}^3F$  term. It will now be shown that the  ${}^3D$  terms are probably responsible, although it is not unambiguously proved.

It is very probable that the  $5d$   ${}^3D$  terms are perturbed in the Sn II spectrum, for both Green and Loring (*loc. cit.*) and Lang (*loc. cit.*) have shown that in addition to the normal doublet system having  ${}^3p$  as its basic configuration, there exists a parallel  ${}^3p^2$  system of doublets and quartets. It is seen from fig. 1 that the  $5d$   ${}^3D$  terms of these two systems lie fairly close to each other. Russell and Shenstone† have shown that it is just such terms, namely those with the same  $n, l, J$  values, but arising from different electron configurations, that perturb very strongly, even if a large interval separates them. The Pb II spectrum is very similar in structure, and the two sets of  ${}^3D$  terms are almost identically situated in both spectra. It is known that there is a strong perturbation between these terms in Pb II so that the same will be likely in Sn II. These perturbed terms in Pb II show abnormally wide even isotope displacement‡. It is therefore to be expected that even isotope displacement should occur in both the  $5d$   ${}^3D_{5/2}$  and the  $5d$   ${}^3D_{3/2}$  terms of Sn II, and as shown previously, the experimental evidence supports this.

The abnormally large value of the  $5d$   ${}^3D_{5/2}$  interval factor is also very probably a result of the same perturbation that produces the isotope displacement. It was shown by Schüller and Jones§ that when fine structure perturbations occur between two terms there is an apparent repulsion between levels of the same  $F$  value. The repulsions are proportional to  $F$  so that if all the levels of a term are perturbed (which is the case when the  $J$  values of the two perturbing terms are the same) the interval rule still holds in the perturbed structures. The perturbation may either close or open up a structure, depending on the relative positions of the perturbing levels. Jones|| also pointed out that the strongly perturbed  ${}^1P_1$  term in Hg I and the  $6d$   ${}^3D$  terms of Pb II are associated with a very large even isotope displacement. Taking all the evidence into consideration, it suggests that the explanation of the abnormal  ${}^3D$  structures is probably to be sought in perturbations. In this respect it may be noticed that the even isotope line in  $5d$   ${}^3D_{5/2}-4f$   ${}^3F$  is not exactly at the calculated centre of gravity of the fine structure multiplet, but is dis-

\* An analogous case exists in the very similar Pb II spectrum.

† 'Phys. Rev.', vol 39, p 415 (1932).

‡ Schüller and Jones, 'Z. Physik,' vol 75, p 563 (1932).

§ 'Z. Physik,' vol 77, p 801 (1932).

|| 'Proc. Phys. Soc.', vol 45, p 501 (1933).

placed to the violet by 10 units, which, although small, is apparently real. Such a displacement would arise if a perturbation took place, as can easily be shown.

### *Nuclear Magnetic Moment*

Recent theoretical developments due to Goudsmit\* make it possible to calculate the nuclear magnetic moment, for a one-electron spectrum, with a high degree of certainty, if the necessary fine structures are known. The values of the nuclear magnetic moments are just as important as the values of the mechanical moments, for although the latter are quantized, the former are not, showing that although the nuclear particles have half integral mechanical spins, these spins are associated with different magnetic moments for different kinds of particles. Goudsmit has shown that, if  $g(I)$  is the ratio of the magnetic to the mechanical moment of a nucleus, the former being expressed in Bohr magnetons, the following relationship holds for the structure in an  $s \ ^2S_{1/2}$  term:

$$g(I) = \frac{3\alpha \ n_0^3 \ 1838}{8 \ R\alpha^2 \ Z_i Z_o^2 \ \kappa(\frac{1}{2}, Z_i)}$$

The symbols have the following values for the present term under consideration —

$\alpha$  is the interval factor of the term and is equal to  $0.207 \text{ cm.}^{-1}$

$n_0 = 2.685$  is the Rydberg denominator of the term which in this case is the  $6s \ ^2S_{1/2}$  term

$R = 109,737$  is the Rydberg constant

$Z_i$  is the effective nuclear charge in the inner portion of the penetrating orbit, and as the fine structure interaction comes from the region very close to the nucleus,  $Z_i = Z$  where  $Z$  is the atomic number. For tin this is 50.

$Z_o$  is the charge in the outer portion of the orbit and since the spectrum is that of Sn II, equals 2.

$\kappa(\frac{1}{2}, Z_i)$  is a calculated relativity correction depending on the value of  $Z$  and  $J$  and is in this case 1.30.

$\alpha^2$  is the spectroscopic doublet constant  $= 5.305 \times 10^{-5}$

On substituting these values into the above equation, the value obtained for  $g(I)$  is  $-1.78$ . Alternatively, the magnetic moment is  $-0.89$  Bohr proton magnetons or  $1/2060$  that of the electron.

\* 'Phys. Rev.', vol. 43, p. 636 (1933)

*Fine Structure in the Sn I Spectrum.*

When the pressure of helium in the tube was reduced, the Sn I line 5631.9 ( $2p\ ^1S_0-3s\ ^3P_1$ ) came up very strongly. The structure of this line is as follows --

$$\begin{array}{ccccccc} + & 158 & 0 & 82 & - \\ & (1) & (10) & (2) & \end{array}$$

As in the Sn II lines, the central strong sharp component is due to the even isotopes and the weaker components to the odd. The analysis of this line is

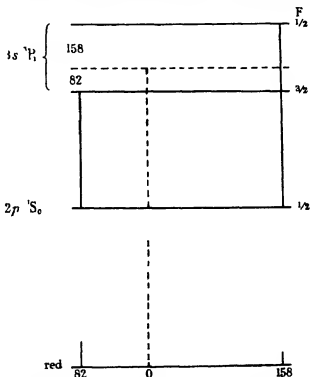


FIG. 5.—Fine structure of the Sn I line  $\lambda$  5631.9 ( $2p\ ^1S_0-3s\ ^3P_1$ )

of particular value, since the lower term has a  $J = 0$  and therefore remains single, no matter what the value of the spin ( $2J + 1 = 1$ ). The two satellites therefore give the structure of the upper term, and as the  $J$  value of this term is 1, the doublet fine structure proves that the spin is  $\frac{1}{2}$ , for any other value would result in a triplet structure. The analysis of the line is shown in fig. 5. The bigger  $F$  level lies deeper, again proving that the magnetic moment is negative. The even isotope line falls extremely close to the calculated optical

centre of gravity of the pattern. The  $F$  values being  $\frac{1}{2}$  and  $3/2$ , the quantum weights of the levels are in the ratio 1 : 2, so that the centre of gravity is 80 units from the stronger component. The measurements give it at 82, which is extremely good agreement.

The interval factor for the  $3s\ ^3P_1$  term of Sn I is 160. It may therefore be taken as definitely proved that the nuclear spin of tin is  $-\frac{1}{2}$ , for this value is obtained from independent analysis of Sn I and Sn II lines. This is true for the two stronger odd isotopes 117 and 119. It is usually considered that as the even isotopes show no structure, this is due to a zero spin, but it is well to point out the no structure is produced when the nuclear magnetic moment is small even when associated with a mechanical moment. It can therefore only be concluded with certainty that the nuclear magnetic moments of 116, 118, 120, 122, 124 are very small, possibly zero. Unfortunately nothing can be said about the weak tin isotopes 115 and 121. These are isobaric with indium and antimony respectively and a comparison of isobar spins would have been of value. Cadmium is the only other atom in which both the odd isotopes are known to possess negative spins. Table IV is of interest, showing how abruptly the spins change from small negative to large positive values, although the atomic weights form a regular series.

Table IV

Atom	Cd	Cd	In	Sn	Sn	Sb	Sb
Isotope	111	113	115	117	119	121	123
Spin	$-1/2$	$-1/2$	$9/2$	$-1/2$	$-1/2$	$5/2$	$7/2$

I wish to take this opportunity of expressing my thanks to Professor A. Fowler, F.R.S., for his continual assistance and encouragement during this work, and for the excellent facilities he has afforded me in his laboratory. I also wish to thank Professor W. E. Curtis, of Newcastle, for the loan of a pair of interferometer plates.

#### Summary

The fine structures of the visible lines of the Sn II spectrum have been analysed by means of a silvered Fabry-Pérot interferometer. The structures of eight terms have been observed, and it is shown that the nuclear spin of the two main odd isotopes 117 and 119 is  $-\frac{1}{2}$ . The nuclear magnetic moment for these isotopes has been calculated and  $g(I)$  found to be  $-1.78$ , giving a magnetic moment  $1/2060$  that of the electron.

Even isotope displacement is found in some lines, and it is shown that this is probably due to perturbations taking place between the  $^2D$  terms arising from the  $s^2 d$  and the  $s p^2$  electron configurations, as in the Pb II spectrum.

The fine structure of the  $2p \ ^1S_0-3s \ ^3P_1$  line of the Sn I spectrum is reported and analysed, and from this a nuclear spin of  $-\frac{1}{2}$  is also deduced, thereby confirming the value obtained from Sn II.

Attention is drawn to the very abrupt changes which take place in the signs and values of the nuclear spins in the series of odd isotopes between 111 and 123.

### *Hyperfine Structure in the Arc Spectrum of Xenon*

By E. GWYNNE JONES, Beit Scientific Research Fellow, Imperial College of Science, London

(Communicated by A. Fowler, F.R.S.—Received November 22, 1933)

The spectrum of neutral xenon (Xe-I) was first analysed by Meggers, de Brun and Humphreys\* in 1929. One of the objects of their investigation was to determine whether any of the lines of xenon would be suitable for consideration as wave-length standards to replace the  $\lambda 6438$  of cadmium, which is considered† to be unsuitable in many respects. The analysis showed that the strongest lines in the region  $\lambda\lambda 4200$  to  $8800$  are  $s-p$  transitions and are therefore theoretically suitable as standards, in the event of their not possessing hyperfine structures, since they are not subject to Stark displacements in the electric field of the discharge‡. It is thus important that the spectrum should be thoroughly examined, with apparatus of the highest resolving power, for hyperfine structure.

#### *Apparatus*

Xenon, on account of its high atomic weight, emits very sharp lines in ordinary light-sources, so that no great precautions were necessary beyond care that lines involving metastable terms (in particular  $1s_2$ ) should not be produced under conditions that would permit self-absorption to occur.

The lines were excited, by means of a transformer supplying about 10,000 volts A.C., in a Pyrex discharge tube, of H-shape, so that both the "end-on"

\* 'Bur. Stand. J. Res.', vol. 3, p. 731 (1929).

† '7th Conf. Gen. Poids et Mesures', p. 52 (1927).

‡ Harkness and Heard, 'Proc. Roy. Soc. A', vol. 139, p. 416 (1933).

and "side on" positions could be used. The lower ends of the tube, which contained the aluminium electrodes, were much lengthened in order to keep the capillary free from sputtered material, as xenon causes very excessive electrode sputtering. Currents ranging from 1/10 to 5 mA were employed—only the weakest excitations being used for lines ending on metastable levels—with exposure times varying from  $\frac{1}{4}$  minute to  $7\frac{1}{2}$  hours.

Fabry-Pérot étalons, mounted in the "parallel beam" of a large-aperture spectrograph, provided the necessary interferometric apparatus. The étalons were of the fixed separation type, with various thicknesses from 5 to 25 mm, the plates being silvered by cathodic sputtering in argon to suitable densities for the different spectral regions. The spectra were recorded on Ilford Monarch, hypersensitive panchromatic or infra-red plates according to the region examined. It was necessary to "sensitize" the infra-red plates with ammonia for all but the strongest lines. About 85 photographs, obtained with various étalon separations, were retained for measurement.

### *Experimental Results*

It was established that the majority of the lines investigated possess fairly complex structures. A few, notably some of those involving the term  $1s_4$ , have no appreciable structures but are slightly broadened, the analysis of other line structures shows that this is due to the small structure of the term  $1s_4$ , which is only recognizable when in combination with another term possessing a greater structure. In general, the line patterns consist of an intense central component, together with a number of fainter lines whose total intensity is roughly the same as that of the central component. In view of Aston's\* abundance ratios and of all other previously analysed hyperfine structures of elements with "odd" and "even" isotopes, it is clear that this strong central component represents the unresolved even isotopes (mass numbers 124, 126, 128, 130, 132, 134, and 136, totalling 52% of the whole) and that the outer, less intense, components are due to the action of nuclear moments in the odd isotopes (129 and 131, respectively 27.1% and 20.7%).

The problem is now to analyse these fainter components into a scheme that will relate the observed structures to definite terms and to assign nuclear moments to account for these structures. It has been found possible† to allocate the line structures to the various terms, to assign the nuclear moment

\* 'Proc Roy Soc,' A, vol 126, p 511 (1930)

† Cf E. G Jones, 'Nature,' vol 132, p. 781 (1933)



$I = \frac{1}{2}$  to  $\text{Xe}_{130}$  and to assert that for  $\text{Xe}_{131}$ ,  $I > 3/2$ . Owing to the fact that the less abundant odd isotope has the larger nuclear spin, it has not been possible to obtain the complete patterns since, for an atom with a nuclear moment greater than  $\frac{1}{2}$ , these contain numerous faint components which are difficult to recognize, particularly when the structures are comparatively narrow. Nevertheless, the analysis can be carried out with certainty for  $\text{Xe}_{130}$  ( $I = \frac{1}{2}$ ), and the term separations for one odd isotope suffice to give a complete survey of the hyperfine structures occurring in the spectrum. For terms of given  $J$  value, the  $\text{Xe}_{131}$  separations will always bear a constant ratio to those of  $\text{Xe}_{130}$ , and thus do not add to the knowledge of the coupling between the nucleus and the valency electrons.

In a very brief note (in which no details are given) Kopfermann\* states that he has also analysed a number of xenon line structures and assigned the nuclear moments  $I_{130} = \frac{1}{2}$  and  $I_{131} > 3/2$ . The two independent analyses thus lead to the same result.

Convincing proof of the correctness of the above interpretation is obtained when the various xenon line structures are compared with those of corresponding mercury lines: a striking similarity is at once evident—the xenon line structures are usually perfect mirror images, on a smaller scale, of the mercury structures. It has been shown†‡ that, for mercury, the even isotopes have zero nuclear spins ( $I = 0$ ), while for the more abundant odd isotope,  $\text{Hg}_{199}$ ,  $I = \frac{1}{2}$  and for the other,  $\text{Hg}_{201}$ ,  $I = 3/2$ . Thus a complete analogy between the structures is to be expected.

### Notation

The term notation adopted by Humphreys and Meggers§ is retained in the present work, but the appropriate  $J$ -value is added as a superscript to the term symbols.

### The Analysis of the Line Structures

The observed line structures and the analysis for  $\text{Xe}_{130}$  are shown in the figs 1 to 9, in which the positions of the components are given in units of  $10^{-3} \text{ cm}^{-1}$ . The intensities of the various components, according to a rough visual estimate, are proportional to the heights of the lines and the theoretical

\* 'Naturwiss.', vol. 39, p. 704 (1933).

† Schüller and Keyston, 'Z. Physik,' vol. 72, p. 423 (1931).

‡ Schüller and Jones, 'Z. Physik,' vol. 74, p. 631 (1932), vol. 77, p. 801 (1932).

§ 'Bur. Stand. J. Res.', vol. 10, p. 139 (1933).

intensities—calculated from the appropriate intensity rules\* and Aston's abundance ratios—are found in the term schemes. The components assigned to  $\text{Xe}_{139}$  are indicated by capital letters, and those assigned to  $\text{Xe}_{131}$  by small letters.

The structure of the line  $\lambda 7967 (1s_3^0-3p_7^1)$  is of particular importance, since the appearance of four components—apart from the central one—proves that the two odd isotopes have different nuclear moments. The term  $1s_3^0$  is characterized by  $J=0$  and thus, in the absence of isotopic displacements, can have no hyperfine structure, hence, the term  $3p_7^1$  has at least four hyperfine sub-levels arising from the odd isotopes. Isotopic displacements are

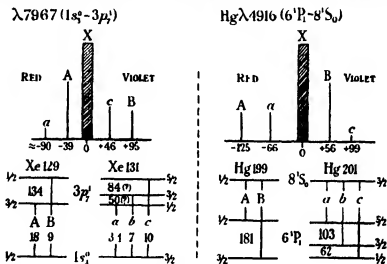


FIG. 1

ruled out by the sharpness of the central component, X, in all lines. Now,  $3p_7^1$  has  $J=1$  and thus cannot have more than a three-fold hyperfine structure unless isotopes with more than one nuclear moment are involved. In view of the fact that  $\text{Xe}_{139}$  is the more abundant odd isotope, the strongest components in the line patterns will belong to it, accordingly in  $\lambda 7967$ , "A" and "B", fig. 1, are assigned to  $\text{Xe}_{139}$ . Their relative intensities ( $\approx 2:1$ ) and positions with respect to the "null-line" indicate that  $I_{139} = \frac{1}{2}$ , as otherwise another component (approximately half the intensity of B) would be required on the "violet" side and also the centre of gravity of this new pattern would no longer coincide with "X". The components "a" and "b" are assigned to  $\text{Xe}_{131}$ , but a third component, between a and X, is required in

\* Güttinger and Pauli, 'Z. Physik,' vol. 67, p. 743 (1931)

order that the centre of gravity of  $Xe_{129}$  should coincide with X. The complete analysis is presented in fig 1, which also contains the structure and analysis of Hg  $\lambda$  4916, likewise a  $J = 0 \rightarrow 1$  transition, and therefore completely analogous to Xe  $\lambda$  7967.

The lines  $\lambda$  7887 ( $1s_2^1 - 2p_1^0$ ) and  $\lambda$  7642 ( $1s_2^0 - 2p_2^1$ ) have structures that may be similarly analysed, fig 2. These two lines are not so favourable as  $\lambda$  7967 for the observation of faint components.

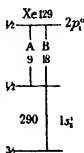
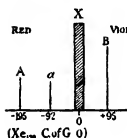
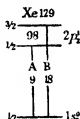
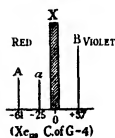
 $\lambda$  7887 ( $1s_2^1 - 2p_1^0$ ) $\lambda$  7642 ( $1s_2^0 - 2p_2^1$ )

FIG 2

The structures of the lines  $\lambda$  8409 ( $1s_2^2 - 2p_1^1$ ) and  $\lambda$  4792 ( $1s_2^2 - 3p_{10}^1$ ) need special comment. They are both exactly analogous to the mercury line  $\lambda$  5769, this agreement is very satisfactory since, in all these cases, both upper and lower terms possess structures and, as a result, the line patterns are rather complex, figs 3 and 4.

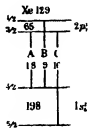
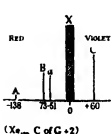
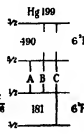
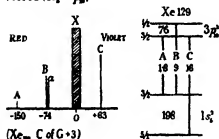
 $\lambda$  8409 ( $1s_2^2 - 2p_1^1$ )Hg  $\lambda$  5769 ( $6^1P_1 - 6^1D_2$ )

FIG 3

The structures of the terms  $1s_2^2$  and  $2p_2^1$  being now determined, it is possible to calculate the positions of the strongest satellites of  $Xe_{129}$  for  $\lambda$  4500 ( $1s_2^2 - 2p_2^1$ ). The agreement with experiment is very close. The structure of the

term  $1s_3^2$  permits the analysis of the lines  $\lambda\lambda$  4203, 4524, 8231, 8819, 4624, 4671, 4697, and 4792, from which the  $Xe_{129}$ -structures can be derived for the terms  $4Y$ ,  $2p_3^2$ ,  $2p_3^1$ ,  $2p_3^0$ ,  $3p_4^2$ ,  $3p_4^1$ ,  $3p_4^0$ , and  $3p_{10}^1$

$\lambda 4792 (1s_3^2 - 3p_4^1)$



$\lambda 4500 (1s_3^2 - 2p_4^1)$

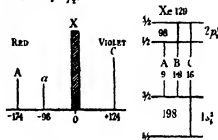
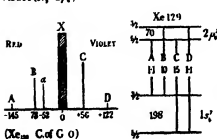


FIG 4

The lines  $\lambda$  4734 ( $1s_3^2 - 2p_3^2$ ) and  $\lambda$  4524 ( $1s_3^2 - 2p_3^2$ ) give the structures of  $2p_3^2$  and the important term  $1s_4^1$ . The former term structure is again checked by the line  $\lambda$  8346 ( $1s_3^2 - 2p_3^2$ ), which also shows two of the strongest  $Xe_{131}$  components,  $a$  ( $F = 5/2 \rightarrow 3/2$ ) and  $b$  ( $F = 5/2 \rightarrow 5/2$ ), indicating that the total separation of the  $Xe_{131}$ -hyperfine structure is about  $0.235 \text{ cm}^{-1}$  for  $1s_3^2$ . The separation assigned to the term  $2p_4^1$  is only approximate, as it is estimated from the fact that  $\lambda$  4690 ( $1s_3^2 - 2p_4^1$ ) possesses no marked structure—showing that the separations of the terms  $1s_3^2$  and  $2p_4^1$  are nearly equal. This estimated structure is in qualitative agreement with the incompletely observed hyperfine structure of the line  $\lambda$  4916 ( $1s_4^1 - 2p_4^1$ ).

$\lambda 8231 (1s_4^1 - 2p_4^1)$



$\lambda 4624 (1s_3^2 - 3p_4^1)$

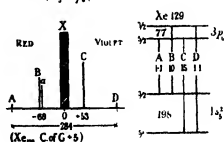


FIG 5

The total separations of the terms of  $Xe_{131}$  appear to be slightly less than those of  $Xe_{129}$ , and are always "inverted" with respect to those of the lighter isotope, proving that the nuclear magnetic moments are of opposite sign. It

may also be shown, by the construction of tentative schemes for  $Xe_{131}$ , that no additional intense components are to be expected in observable positions for any of the lines. Thus the proposed analysis explains all the observed structures of the 20 principal Xe-I lines

$\lambda 4697(1s_2^2 - 3p_2^2)$

$\lambda 4203(1s_2^2 - 4Y^1)$

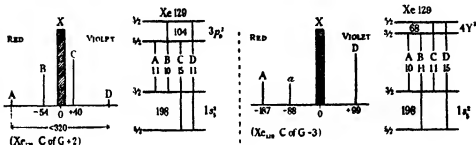


FIG. 6

### Discussion of Results

The hyperfine separations (for the isotope  $Xe_{131}$ ) of the terms of the Xe-I spectrum, as derived from the preceding analysis of the line structures, are collected in Table I, which also shows the lines that were used in the determination of the particular term structure. It is noteworthy that, in all terms

$\lambda 8819(1s_2^2 - 2p_2^2)$

$\lambda 4671(1s_2^2 - 3p_2^2)$

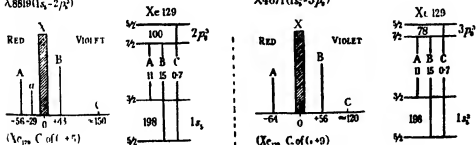


FIG. 7.

except  $2p_2$  and  $4Y$ , the hyperfine level of greater  $F$ -value is lower in the energy diagram (i.e. the hyperfine levels of  $Xe_{131}$  are usually "inverted")

No trace of an isotopic displacement was observed in any of the lines, the component (X) due to the even isotopes is always quite sharp

Table I—Hyperfine separations of the terms of  $\text{Xe}_{139}$  ( $I = \frac{1}{2}$ )

Term	$I$ value	Hyperfine separation	Landé interval factor	Lines involved
		$\text{cm}^{-1}$	$\text{cm}^{-1}$	$\lambda$
$1s_{\frac{1}{2}}^1$	1	-0 290	-0 193	7887, 8346
$1s_{\frac{3}{2}}^1$	0	0 0	0 0	7967, 7642
$1s_{\frac{1}{2}}^1$	1	-0 057	-0 038	4734
$1s_{\frac{3}{2}}^1$	2	-0 198	-0 079	8409, 4792, 4500, 8231, 4624 4203, 8819, 4671, 4524, 4697
$2p_{\frac{1}{2}}^1$	0	0 0	0 0	7887, 4582
$2p_{\frac{3}{2}}^1$	1	+0 008	+0 065	7642, 4500
$2p_{\frac{3}{2}}^1$	2	-0 250	-0 100	4734, 4524, 8346
$2p_{\frac{1}{2}}^1$	1	-0 220	-0 147	(4616), (4690)
$2p_{\frac{3}{2}}^1$	0	0 0	0 0	8280
$2p_{\frac{3}{2}}^1$	2	-0 070	-0 028	8231
$2p_{\frac{1}{2}}^1$	1	-0 065	-0 043	8409
$2p_{\frac{3}{2}}^1$	3	-0 100	-0 029	8819
$3p_{\frac{1}{2}}^1$	0	0 0	0 0	4807
$3p_{\frac{3}{2}}^1$	2	-0 076	-0 030	4624
$3p_{\frac{1}{2}}^1$	1	-0 134	-0 089	7967
$3p_{\frac{3}{2}}^1$	1	-0 078	-0 022	4671
$3p_{\frac{1}{2}}^1$	2	-0 104	-0 042	4697
$3p_{\frac{3}{2}}^1$	1	-0 076	-0 051	4792
$4Y$	2	+0 068	+0 027	4203

The '-' sign indicates that the hyperfine levels are inverted

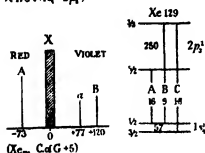
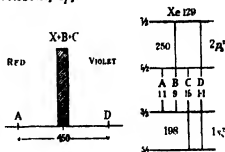
 $\lambda 4734 (1s_{\frac{1}{2}}^1 - 2p_{\frac{3}{2}}^1)$  $\lambda 4524 (1s_{\frac{1}{2}}^1 - 2p_{\frac{3}{2}}^1)$ 

FIG. 8

The  $mp_1$ - and  $mp_3$ -series show the existence of selective perturbations in the hyperfine structures of the members with  $m = 3$ , each perturbation is perceptible as an anomalous increase in the hyperfine separation of the term when compared with the lowest member. The quantum defects of the terms are also anomalous and likewise indicate the presence of perturbations at  $m = 3$ . Exactly similar perturbations of hyperfine structures have been described\* in the spectra Hg-I and Pb-II

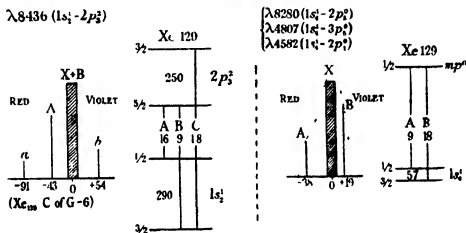
\* E. G. Jones, 'Proc. Phys. Soc.', vol. 45, p. 501 (1933)

In conclusion, the writer wishes to express his thanks to Professor A. Fowler, F.R.S., in whose laboratories the work was carried out, for his kindly interest and encouragement.

### Summary

(1) All the important lines in the region  $\lambda\lambda$  4200 to 8900 of the spectrum of neutral xenon have been investigated, with Fabry-Pérot étalons, for hyperfine structure. It was found that 16 lines were complex, so that it may be concluded that the lines of Xe-I are not suitable as wave-length standards.

$\lambda 8436 (1s_1' - 2p_2')$



(2) The observed line structures have been analysed and the corresponding term structures determined. These structures are explicable if the nuclear moments  $I = 0$  be assigned to the even isotopes,  $I = \frac{1}{2}$  to  $\text{Xe}_{130}$  and  $I > \frac{3}{2}$  to  $\text{Xe}_{131}$ . The nuclear magnetic moment of  $\text{Xe}_{131}$  has a sign opposite to that of  $\text{Xe}_{130}$ .

(3) No isotopic displacements were found in any of the lines and the observed intensities of the components in the line patterns indicate that the even mass-numbers represent about 50% of the xenon isotopes.

# *The Scattering of Electrons in Ionizing Collisions with Gas Atoms*

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(Communicated by Lord Rutherford, O.M., F.R.S. — Received November 23, 1933)

## *Introduction*

When electrons lose energy by colliding with gas atoms, they may do so either in definite amounts by exciting the atom to a discrete level, or by ionizing the atom, in which case the electrons may lose any amount of their energy over and above the minimum required for ionization. The first process has now been studied in several aspects by a number of investigators, in particular, the authors\* have dealt with the angular distribution of the scattered electrons.

The study of ionizing collisions, however, has so far been confined largely to the measurement of collision cross-sections,† while only slight attention has been paid to the angular distribution and the energy distribution of the scattered electrons.‡ The most valuable information concerning the collision process is to be derived from a detailed study of the angular distribution of the scattered electrons, and such an investigation is described in the present paper.

Little work has so far been done on the angular distribution of electrons scattered in ionizing collisions. Tate and Palmer (*loc. cit.*) have carried out a few measurements in mercury vapour, in which the collected electrons were separated into groups having a large energy spread. In view of the results of the present paper, however, it is clear that reasonably good velocity resolution of the collected electrons is necessary in order to observe the interesting phenomena which occur. Hughes and McMillen (*loc. cit.*) have given some scattering curves in argon and hydrogen for collected electrons which had only a few volts energy. The results they have obtained will be discussed later.

The present paper describes experiments on the angular distribution of electrons which have lost different amounts of energy over and above the

\* 'Proc. Roy. Soc., A,' vol. 142, pp. 320, 647 (1933).

† Bleakney, 'Phys. Rev.,' vol. 35, p. 139 (1930), Smith, 'Phys. Rev.,' vol. 36, p. 1293 (1930), Tate and Smith, 'Phys. Rev.,' vol. 39, p. 270 (1931).

‡ Tate and Palmer, 'Phys. Rev.,' vol. 40, p. 731 (1932), Hughes and McMillen, 'Phys. Rev.,' vol. 39, p. 585 (1932), vol. 41, p. 39 (1932).



amount required for ionization, and the investigation includes measurements on electrons which leave the atom with certain definite velocities. Electrons with different incident energies have been used, and measurements have been carried out in hydrogen, helium, nitrogen, methane, neon, argon, and mercury vapour.

### Method

The apparatus used in the present work has already been described in detail in a previous paper\*. The essential parts are a collision chamber containing the gas at a low pressure, and a highly evacuated chamber containing an electrostatic analyser. Scattered electrons entering the analyser are subjected to a velocity analysis, and the number of electrons with the desired energy is measured as a function of the angle of scattering by rotating the electron gun.

The precautions taken in obtaining measurements have been discussed elsewhere† in connection with the discrete losses, *i.e.*, losses corresponding to excitation of the atom to a discrete level. With ionization losses there are no peaks in the velocity distribution of the collected electrons such as occur for the main discrete losses, and one must be certain that the measured effects are not partly due to 'background'. Consideration shows that background may arise from two causes: (a) stray scattering in the collision chamber, most probably from the slits and walls, (b) stray electrons in the analyser reaching the collector with energies different from that which the analyser is adjusted to focus. The following tests indicated that such background effects were quite unimportant in the present work—

- (i) When there was no gas in the collision chamber, the collected current was only a small fraction of that when gas was present.
- (ii) The intensity of collected electrons with energies between those of the elastically and inelastically scattered electrons was always small compared with the intensity of the electrons which had made ionizing collisions.
- (iii) For investigations on electrons which retained only a few volts energy after collision, it was convenient to accelerate the scattered electrons up to 30 volts energy before they entered the analyser. Under these circumstances the intensity of the collected current fell abruptly to a very small value when the analyser was adjusted to focus electrons with energies slightly less than 30 volts.

\* 'Proc Roy Soc,' A, vol 142, p 320 (1933)

† 'Proc Roy Soc,' A, vol 138, p 229 (1932)

Test (i) shows that background due to stray scattering in the collision chamber was small compared with the scattering by the gas, and tests (ii) and (iii) showed the absence of appreciable background due to stray electrons getting around the analyser and entering the collector. Further evidence of the reliability of the results was obtained from the fact that the curves were closely reproducible at different gas pressures.

### Results

(a) *Argon and Mercury Vapour* - The ionization potential of argon is 15.7 volts, and angular distributions were measured for electrons which had lost this or a greater amount of energy in the scattering process. In fig. 1\* curves are given† for incident electrons of 100, 60, 40, 30 and 20-volts energy. The 100-volt curves are for electrons which have lost 15.7, 25, 40, 65, 75 and 95-volts energy in the collision. Similarly, curves for various losses greater than the ionization potential are given for the other incident velocities. The ionization potential of mercury vapour is 10.4 volts, and in fig. 2 curves are given for incident electrons of 60, 40, 30, 20 and 14-volts energy.

These curves exhibit some features which are also present in the curves for the other gases studied. These features are -

- (i) A close similarity between the curves for the ionization loss and the corresponding curves for the main discrete loss, though at small angles the former curves are less steep than the latter, this last effect is more marked at the lower incident voltages.
- (ii) For incident electrons of any particular energy, the curves for electrons which have lost the larger amounts of energy are flatter than those which have lost smaller amounts in the collision. Further, the change in steepness is particularly rapid within a range of a few volts near the voltage at which the scattered electrons have half the energy of the incident electrons, this effect, however, is not so marked in the curves for argon, and is even less so for mercury vapour.

Turning now to the curves for argon and mercury vapour in particular, we observe further features of interest. Diffraction effects, in the form of a maximum and a rise at large angles, are present in the curves even when the collected electrons have only a few volts energy. Moreover, the curves for

\* In this and subsequent figures, all the curves are drawn to arbitrary scales.

† The zeros of the successive curves are given on the right-hand side.

electrons which have only 5 volts energy after the collision are markedly similar for all incident electron energies. Such curves for argon are different from the corresponding ones for mercury vapour, in which the rise at small angles is more pronounced.

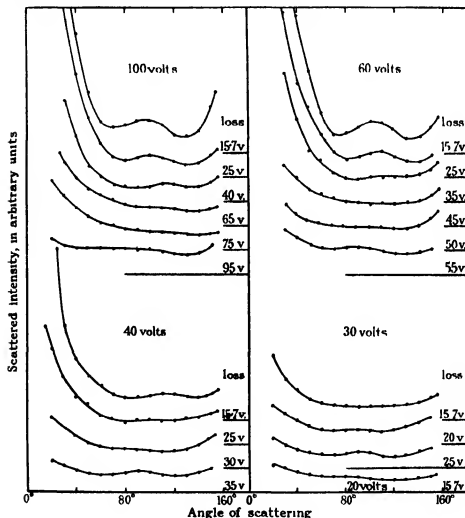


FIG 1—Scattering in argon

In the curves for argon, as the energy lost by the electron increases, the maximum tends to move out to a slightly larger angle and then moves in to a smaller angle, while its prominence decreases and then increases again slightly. Similar but less pronounced effects are present in the curves for mercury vapour.

Hughes and McMillen (*loc cit*) have given curves for the scattering of electrons in argon and hydrogen for collected electrons with energies of 1, 3, 5.5 and 8-volts energy. Most of these curves show very sharp maxima at large angles, the curves being otherwise fairly flat. Our attempts to verify

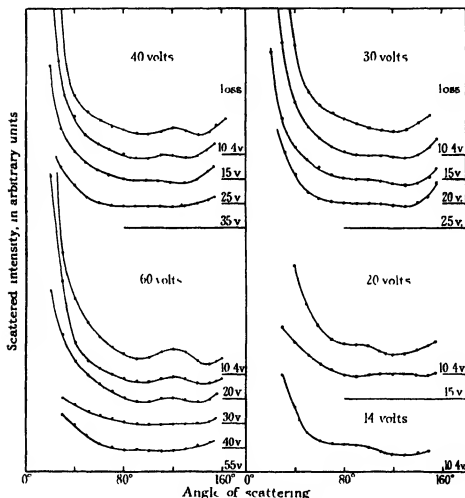


FIG. 2—Scattering in mercury vapour

the presence of such maxima were unsuccessful, however, too much importance should not be attached to this result, since in some experiments when studying collected electrons of very low energy, there was a background due to slow electrons ejected from the walls of the scattering chamber. Background effects were, of course, quite small for all the curves given in this paper.

(b) *Helium*—In fig 3 scattering curves are given for electrons with incident energies of 200, 100, 60 and 40 volts, and for various losses greater than the ionization potential of 24.5 volts. At large angles all the curves are quite flat, and in general they show the characteristics (i) and (ii) already mentioned above. In addition to these characteristics some of the curves show a kink, a feature which is seen later to be of some significance.

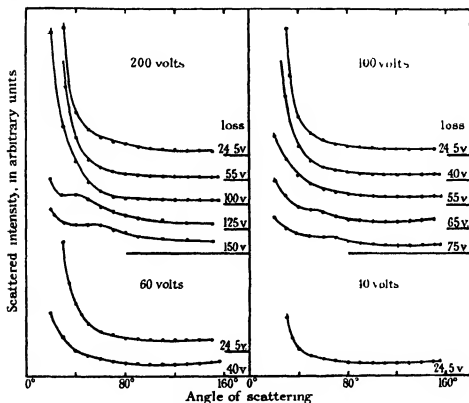
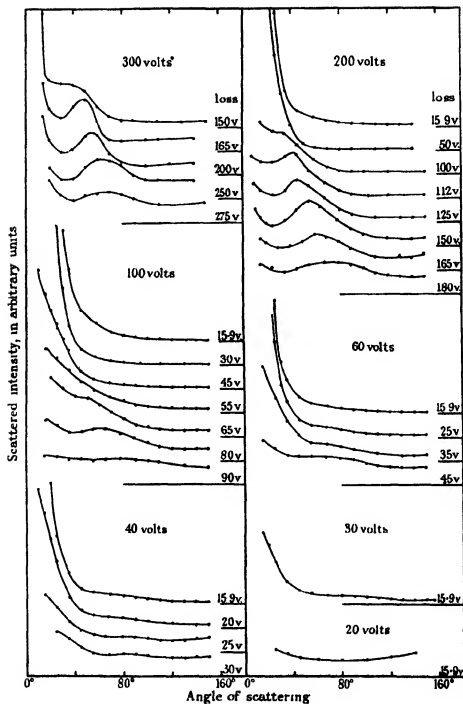


FIG 3—Scattering in helium

(c) *Hydrogen*—In fig 4 curves are given for the scattering of 300, 200, 100, 60, 40, 30 and 20-volt electrons in hydrogen, the ionization potential of which is 15.9 volts. At the higher voltages, many of the curves are seen to exhibit a striking and unusual feature in the occurrence of a prominent maximum, which moves regularly as the energy loss changes. As the amount of energy lost by the electron decreases, the maximum moves into smaller angles, until it is finally lost in the increased steepness of the curves. This type of maximum is not present in the inelastic curves for the main discrete loss, and it will be



clear that it is of quite a different nature to that observed at large angles in the curves for argon and mercury vapour. At the lower incident voltages the effect is still present although much less pronounced, the maximum being reduced to a slight undulation.

(d) *Nitrogen, Methane and Neon* — Since the very interesting feature observed in hydrogen was also noticeable to a slight extent in helium, it was clearly of interest to investigate other light gases at higher voltages where the effect would be more pronounced. Measurements were therefore obtained for the scattering of 300, 200 and 100-volt electrons in nitrogen, and for 200-volt

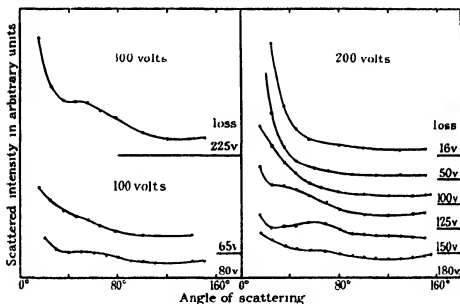


FIG. 5—Scattering in nitrogen

electrons in methane and neon the resulting curves are shown in figs 5 and 6.

The effect observed in hydrogen and helium is also present to a slight degree in these curves, although for neon, the heaviest of the three gases, it has almost disappeared. The main point of interest is that the position of the maximum in corresponding curves for the same incident voltage is independent of the nature of the gas.

#### Discussion

The chief interest attached to the scattering of electrons by the heavier atoms lies in the occurrence of diffraction maxima and minima at large angles

The presence of these diffraction effects is due to the distortion of the incident and outgoing electron waves by the field of the atom. In a previous paper\* the authors have given a qualitative explanation of inelastic scattering in different gases for electrons which have lost a discrete amount of energy. It was shown that when the wave lengths of the incident and outgoing electrons are nearly the same, the curves for inelastic scattering will exhibit a diffraction effect †. For the same reasons it is to be expected that curves for the ionization loss will be similar to the corresponding curves for the discrete loss, provided that the energy of the incident electrons is large compared with the ionization potential. However, when the incident electrons lose an amount of energy

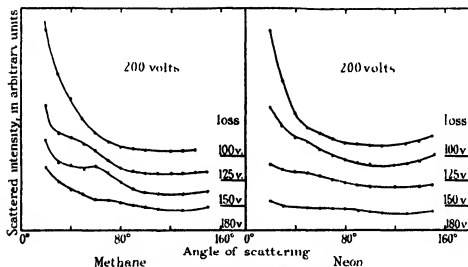


FIG. 6.—Scattering in methane and neon

considerably more than is required to ionize the atom, the wave-lengths of the incident and outgoing electrons are appreciably different, and so the diffraction effect in the curves becomes less prominent. These features are observed in the curves for argon and mercury vapour. Now, when electrons leave the atom with only a very few volts energy, they are moving slowly in the comparatively strong field of the ionized atom, an influence which is independent of, and more important than the energy of the incident electrons. Thus, we expect the angular distributions of electrons leaving the atom with 5-volts

\* 'Proc Roy Soc,' A, vol. 138, p. 469 (1932)

† [Note added in proof, January 9, 1934.—Detailed calculations (to appear shortly) have since been carried out by Massey and Mohr and their results establish this explanation.]



energy (see figs 1 and 2) to be similar for all energies of the incident electron

Let us now turn to the results obtained for the scattering by light elements. In some of the angular distributions a maximum appears which is particularly marked for hydrogen. This maximum becomes less prominent as the energy of the incident electrons decreases, whereas this is not so for the usual diffraction maxima which occur at large angles. Further, the position of the maximum depends only on the energies of the incident and outgoing electrons, and not on the nature of the gas. The effect therefore involves, not the atom, but the atomic electrons, and so considerations of energy and momentum should explain the phenomenon.

Let us therefore consider the collision of an incident and an atomic electron neglecting the effect of the atomic field. Let  $V$  be the energy of the incident electron in volts,  $v, v'$  the energies of the two electrons after the collision,  $\theta, \theta'$  the angle which the directions of the two outgoing electrons make with the incident electron, and  $V_0$  the ionization potential of the atom. The energy and momentum relations then give

$$V^{\frac{1}{2}} = v^{\frac{1}{2}} \cos \theta + v'^{\frac{1}{2}} \cos \theta',$$

$$0 = v^{\frac{1}{2}} \sin \theta - v'^{\frac{1}{2}} \sin \theta',$$

$$V = v + v' + V_0$$

It follows that

$$\cos \theta = \frac{v + \frac{1}{2}V_0}{V^{\frac{1}{2}}v^{\frac{1}{2}}} \quad (1)$$

From this relation we obtain the angle at which electrons with energy  $v$  leave the atom after the collision. We should therefore expect a maximum at an angle  $\theta$  in the angular distribution of collected electrons of energy  $v$ . This maximum will, of course, be superimposed on the angular distribution owing to the scattering by the atomic field. In Table I are compared the observed and calculated angles at which the maximum appears in the various curves for hydrogen, and it is seen that there is remarkably good agreement. This agreement will also hold for other gases, since the calculated position of the maximum depends only slightly on the ionization potential of the atom.

Let us now consider the application of quantum theory to the calculation of the actual form of the curves. Using Born's formula, Massey and Mohr\* and Wetzelt† have carried out calculations for the ionization of the simplest

\* 'Proc Roy Soc.,' A, vol 140, p. 613 (1933)

† 'Phys. Rev.,' vol 44, p 25 (1933)

elements, hydrogen and helium, by electron impact. Wetzel uses the somewhat doubtful approximation of taking plane waves to represent the electron ejected from the atom, while the more exact continuous wave functions were used by Massey and Mohr. Only a few curves were given in their paper owing to the tedious nature of the calculations, and Wetzel does not give curves for hydrogen or for the larger energy losses in helium. Additional curves were therefore

Table I—Calculated and Observed Positions of the Maxima in the Angular Distributions for Hydrogen

V (volts)	$v$ (volts)	$\theta$ calculated (degrees)	$\theta$ observed (degrees)
300	150	42	40
	135	45	48
	100	51	54
	50	63	64
	25	68	68
200	88	43	42
	75	47	44
	50	54	55
	35	59	58
	20	64	72
100	35	41	48
	20	51	60

calculated, using Massey and Mohr's formulae, so as to enable direct comparison with the experimental curves. In fig. 7 theoretical and experimental curves are given for the scattering at small angles in hydrogen and helium, and it is seen that they agree fairly well except in one case. In fig. 8 a number of theoretical curves are given for the scattering of 200-volt electrons in hydrogen and helium. Now the experimental measurements include both electrons scattered by the atom and those ejected from the atom in the collision process, as the two types are experimentally indistinguishable. It would therefore be necessary to calculate the angular distributions for both types and combine them in the appropriate manner for comparison with the experimental curves. It is seen, however, in fig. 8 that the two types of electrons have very similar angular distributions, as might be expected, and so it is usually sufficient to calculate merely the angular distribution of the scattered electrons.

Comparing the theoretical curves in fig. 8 with the corresponding experimental curves for hydrogen and helium in figs. 3 and 4 respectively, we see that there is qualitative agreement in the form of the curves, and that the position of the maximum, when it occurs is also given

correctly. The theory, however, does not give sufficient scattering at large angles. This also occurs when Born's formula is used to calculate the inelastic scattering for the discrete loss, and it has been shown\* that this is due to neglect of the distortion of the incident and outgoing electron waves by the atomic field. Inclusion of this effect for ionizing collisions would, as for the discrete loss, increase the calculated scattering at large angles. The theoretical curves

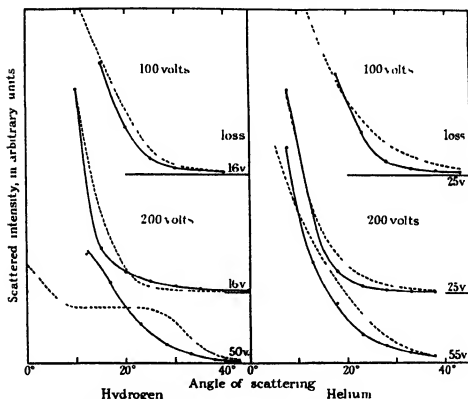


FIG 7 □— Experimental curves, — — — theoretical curves

show a less pronounced and broader maximum for helium than for hydrogen, and inspection of the formula which was used shows that for electrons of the same incident velocity, the maximum becomes much less pronounced for atoms of greater atomic number, the formula also shows that the maximum becomes more prominent at higher velocities. The experimental curves exhibit all these features, as we have already seen

\* Massey and Mohr, 'Proc Roy Soc,' A, vol 139, p. 187 (1933)

One of us (C B O M.) is indebted to the Department of Scientific and Industrial Research for a Senior Research Grant during the latter portion of this work. In conclusion, the authors wish to thank Lord Rutherford and Dr J Chadwick for their interest and encouragement.

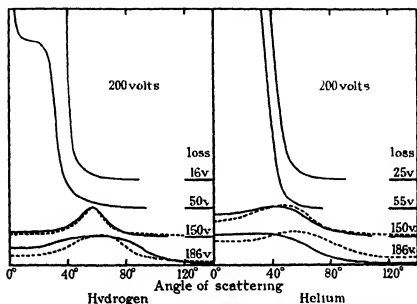


Fig 8—Theoretical curves, — scattered electrons, - - - ejected electrons

### *Summary*

An apparatus, described in a previous paper, has been used for investigations on the angular distribution of the scattering of electrons in gases for ionizing collisions. Measurements were carried out for electrons which had lost various definite amounts of their incident energy over and above that required for ionization. Results have been obtained for hydrogen, helium, nitrogen, methane, neon, argon, and mercury vapour, and curves are given for several incident energies below 300 volts.

The curves obtained exhibit several features which are discussed in detail. An interesting effect is observed in the light gases, particularly in hydrogen, namely, the occurrence of a single pronounced maximum in the angular distribution at certain angles between 30° and 80°. This effect, which is more pronounced at higher velocities, is explained by simple momentum and energy considerations, and curves are obtained with the use of quantum theory which are in fair agreement with the experimental curves.

*An Extension of Southwell's Method of Analysing Experimental Observations in Problems of Elastic Stability*

By H R FISHER, B A.

(Communicated by R V Southwell, F R S—Received December 7, 1933)

1 INTRODUCTION

In a recent paper\* Southwell deals with a well-known type of elastic instability which he exemplifies by considering an axially loaded strut. When a system having such instability is subjected to some type of load  $P$  there corresponds to each one of certain kinematically possible modes of distortion a certain critical load,  $P_1$ , at which this mode becomes unstable. If before loading the system has already a small distortion answering to the mode considered, then the application of  $P$  increases the amplitude of this mode, and hence the representative deflection, in the ratio  $P_1/(P_1 - P)$ . Alternatively it may be said that owing to the application of  $P$  there occurs an *additional deflection*  $\delta$  equal to  $P_1/(P_1 - P)$  times the initial deflection  $\delta_0$ , that is

$$\frac{P_1 - P}{P} \delta = \delta_0,$$

or

$$P_1 v - \delta = \delta_0, \quad (1)$$

if  $v$  be written for  $\delta/P$ .

In most practical cases only the mode which is associated with lowest  $P_1$  (that is, the "fundamental mode") will be important, but it will often be difficult to measure  $\delta$  or  $\delta_0$  without including components arising from other modes of deflection, and then (1), regarded as an equation in  $\delta$  the measured additional deflection, will be only an approximation, which, however, becomes increasingly close as the approach of  $P$  towards  $P_1$  causes the fundamental mode to predominate.

In the paper cited, Southwell pointed out that, in accordance with (1) above, the points obtained by plotting  $\delta$  (the observed deflection) against  $v$  (the deflection divided by the load) may be expected to lie upon a straight line whose slope defines the critical load  $P_1$  (see fig. 1). In the problem of the axially loaded strut  $P$  is the axial load,  $\delta_0$  and  $\delta$  are the initial and additional small transverse deflections at the mid-length, and  $P_1$  is the well-known Euler

\* 'Proc Roy Soc,' A, vol 135, p 601 (1932)

buckling load  $\pi^2 EI / (2l)^2$ , where  $EI$  is the flexural rigidity and  $2l$  the length of the strut \*

Southwell's method of plotting has proved useful in the analysis of experiments on the instability of strips or panels of thin metal sheet subjected to shear † Here  $P$  is the shearing stress, and if (as usually happens) there is

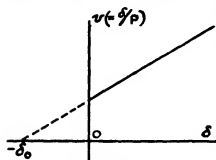


FIG 1—Ideal  $v$   $\delta$  relation

initially any slight deviation from plane-ness whose fundamental component is  $\delta_0$ ,  $\delta$  is the additional depth of the waves or corrugations which are exhibited as the load is increased

11 The following notes develop the idea, suggested by Southwell's example, that the straight-line plotting indicated by (1) might be applicable to the analysis of the mechanical tests which are regularly made on aeroplane spars of

new design The theoretical results may be of interest inasmuch as they illustrate a few elaborations of the method which are apparently of general application

In the tests referred to, a specimen, chosen of such length as to represent an appropriate portion of the whole spar, is subjected to gradually increasing loads approximating in type to those which the spar sustains in service In a

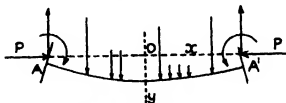


FIG 2—Methods of loading a spar

biplane, or braced monoplane, the loads on a spar (which extends horizontally in the wing from root to tip) consist of compressive axial loads due to the horizontal components of the forces in the external diagonal bracing, combined with transverse loads due to the vertical aerodynamic forces transmitted through the fabric and ribs The types of loading which would accordingly be employed in testing are those indicated in fig 2,  $AA'$  being the centre line of

\* In Southwell's paper (*loc. cit.*) the length of the strut is denoted by  $l$ .

† Gough and Cox, 'Proc Roy Soc,' A, vol 137, p. 145 (1932).

the spar. Actual conditions would usually be symmetrical about the mid-length, but asymmetry will be permitted in what follows, since it introduces little complication. If, in fact, only the mid-length deflection is considered, the anti-symmetrical components of the loads and end eccentricities have no effect.

In spar tests of the standard type the main transverse loads are kept proportional to the end load  $P$  while  $P$  is varied, although constant transverse loads (such, for example, as the weight of the spar) may co-exist. In the present paper both kinds of loading will be taken into account, and it will be shown that in a sufficiently wide range of circumstances an equation of type (1) does theoretically hold with close approximation. The practical advantages of this result are manifest. By fitting a straight line by eye to points plotted on a  $\delta, v$  diagram, a sufficiently accurate value of the flexural rigidity of a spar can be obtained from a large number of experimental readings in a few minutes, and every observation is given weight in arriving at this experimental estimate. Also anomalous behaviour of a tested spar—due, for example, to the limit of proportional elasticity having been exceeded—is immediately recognizable in a departure from straightness of the  $\delta, v$  line. Since  $\delta$  and  $v$  can quite practicably be plotted concurrently with the progress of the test, it is thus possible to obtain early warning of the approach of failure when, as usually, this occurs by yielding or wrinkling of the material at loads well below the critical load  $P_1$ .

1.2 The problem to be considered in this paper can accordingly be enunciated as follows: "By what simple methods is it possible to describe or analyse the behaviour of a uniform beam, initially slightly bowed, which is subjected to the combination of loads shown in fig. 2?" These loads, with the requisite notation, are detailed below.

(i) Compressive loads  $P$  (which are varied during the test) are applied at fixed points in the end cross-sections of the beam. The beam is assumed to be not perfectly straight, and the end thrusts  $P$  are assumed to be applied with some small eccentricity: thus if the line of action of  $P$  be taken as  $Ox$ , with  $O$  at mid-length and  $Oy$  perpendicular as shown, the equation of the unstrained centre line is taken as given by  $y = \eta(x)$ , where  $\eta$  is a function at present unspecified. The end eccentricities are  $\eta(\pm l)$  where  $2l$  is the length of the beam.

(ii) Forces proportional to  $P$  are applied parallel to  $Oy$  at various points along the beam, and couples proportional to  $P$  are applied in the plane  $Oxy$  at the ends. These loads, with the requisite reactions at the ends, are defined

by saying that they produce, in the absence of other loads, a bending moment  $P$  times  $N(x)$  at a distance  $x$  from  $O$

(iii) Constant forces parallel to  $Oy$  are applied at various points along the beam, and constant couples in the plane  $Oxy$  are applied at the ends. These loads, with the requisite reactions at the ends, are defined by saying that they produce, in the absence of other loads, a bending moment  $M(x)$  at a distance  $x$  from  $O$ , and a consequent deflection relative to the ends of amount  $m(x)$ , where

$$-EI \frac{d^2 m}{dx^2} = M, \quad (1A)$$

$EI$  being the flexural rigidity

The problem here presented can, of course, be solved exactly on the basis of the ordinary theory of flexure, the suggested interest of the treatment to be given in this paper centres in the illustration which it affords of principles and methods believed to have applications in other cases of elastic instability. It will be shown that equation (1), regarded as a first approximation, will apply in a wide range of systems, the order of its approximation will be examined, and a second approximation of high accuracy will be suggested.

## 2 THEORY

### *Exact Form of the $v - \delta$ Relation The Function $F(\theta, z)$*

2.1 In the notation of § 1.2, and if  $\delta(x)$  is the additional deflection due to  $P$  and its associated loads—that is, the deflection which ensues on the application of (i) and (ii) after (iii) has already been applied—the bending moment at a distance  $x$  from  $O$ , fig. 2, is

$$Py + M(x) + P N(x),$$

where

$$y = \eta(x) + m(x) + \delta(x),$$

and this bending moment is to be equated to  $EI$  times the excess of the curvature over that implied by  $\eta(x)$ . So we have

$$-EI \frac{d^2}{dx^2} \{\delta(x) + m(x)\} = P \{\eta(x) + m(x) + \delta(x)\} + M(x) + P N(x)$$

Remembering equation (1A) and putting  $P/EI = \mu^2$ , we may write this equation in the form

$$\begin{aligned} \left[ \frac{d^2}{dx^2} + \mu^2 \right] \delta(x) &= -\mu^2 \{\eta(x) + N(x) + m(x)\}, \\ &= -\mu^2 H(x), \quad \text{say} \end{aligned} \quad (2)$$



The deflection  $\delta(x)$  is determined by (2) and by the condition  $\delta = 0$  at  $x = \pm l$ . We observe that there is an exact correspondence between  $\eta$ ,  $N$ , and  $m$ , in that only their sum,  $H$ , affects  $\delta$ . Thus initial bowing, end-eccentricity, and transverse loads and end-couples, whether proportional to  $P$  or constant, all combine to form one total equivalent eccentricity\*.

It is convenient to use instead of  $x$  a co-ordinate  $\theta$  varying linearly from 0 to  $\pi$  along the beam. On introducing the substitutions

$$\theta = (l + x) \frac{\pi}{2l}, \quad D \equiv \frac{d}{d\theta} \equiv \frac{2l}{\pi} \frac{d}{dx},$$

we may write (2) in the form

$$[D^2 + z] \delta = -zH(\theta), \quad (3)$$

where

$$z = \mu^2 \frac{(2l)^2}{\pi^2} = P \frac{(2l)^2}{\pi^2 EI} \quad (4)$$

Now let  $H(\theta)$  be expanded in the range  $(0 < \theta < \pi)$  in the Fourier series

$$H(\theta) = H_1 \sin \theta + H_2 \sin 2\theta + \dots + H_n \sin n\theta + \dots \quad (5)$$

On assuming a similar series for  $\delta$ , substituting in (3) and equating the coefficients of corresponding terms, we deduce that

$$\delta = \frac{z}{1-z} H_1 \sin \theta + \frac{z}{4-z} H_2 \sin 2\theta + \dots + \frac{z}{n^2-z} H_n \sin n\theta + \dots, \quad (6)$$

provided that certain operations on the infinite series are justified. The justification is clear if  $H$  is continuous and zero at the end-points, as was assumed in Southwell's paper. The latter condition is not necessary, however, as will be shown later.

Equation (6) illustrates the opening remarks of this paper. In the expression for  $\delta$ ,  $H_1 \sin \theta$ , the fundamental component of  $H$ , is multiplied by  $z/(1-z)$ , that is, from (4), by  $P/(P_1 - P)$  if (as in the case of the simple strut) we write  $P_1$  for  $\pi^2 EI/4l^2$ , the Euler buckling load. Thus  $z$  measures the end-load in units of  $P_1$ . If we neglect all terms in the series (6) except the first, and

\* We could have foreseen the equivalence of  $N$  and  $\eta$  as regards constant additions, since the same net loading is obtained by decreasing the end-eccentricities and increasing by the same amount the contribution of the end-couples to  $N$ . Nor is it to be supposed that the idea of a general correspondence is new; it is implicit in the various approximate formulae, well known to engineers, which have been given by Perry, Berry, Webb and others for applications to problems of beams loaded axially and transversely.

multiply by  $(1-z)/z$ , we obtain equation (1), except that  $\delta_0$  is replaced by  $H_1 \sin \theta$

The complete equation (6) may be written in the form

$$\frac{1-z}{z} \delta = P_1 v - \delta = H(\theta) \frac{H_1 \sin \theta + \frac{1-z}{n^2-z} H_n \sin n\theta + \dots}{H_1 \sin \theta + \dots + H_n \sin n\theta + \dots} \\ = H(\theta) F_H(\theta, z), \text{ say,} \quad (7)$$

where  $v = \delta/P$  as before. The form of the non-dimensional 'error-function'  $F_H$  as thus defined does not depend upon the magnitude of  $H$ , but only on its distribution along the beam's length.

In order that the graph of  $v$  against  $\delta$  should be a straight line with slope  $1/P_1$ , it would be necessary that  $F_H(\theta, z)$  should be independent of  $z$ . As a rule this is not exactly true, but since the coefficient of  $H_n \sin n\theta$  in the numerator varies from only  $1/n^2$  to 0 when  $z$  varies from 0 to 1, the variation of  $F_H$  will be extremely small if, as usually happens,  $H_1 \neq 0$  and the following few coefficients  $H_n$  are comparatively small. If  $H$  is symmetrical or  $\delta$  is measured at the mid-length only, the second (and all even) terms in the numerator are absent, and the variation is further reduced.\*

#### *Validity of the Series for $\delta$*

2.11 Let  $H$  be continuous, and have limited total fluctuation,† in  $0 \leq \theta \leq \pi$ . Then  $H = \Sigma [H_n \sin n\theta]$  in  $0 < \theta < \pi$ , and

$$H_n \equiv O(1/n) \ddagger \quad (9)$$

Let

$$\delta = \Sigma \left[ \frac{z}{n^2 - z} H_n \sin n\theta \right] \quad (6) \text{ bis}$$

The convergence of this series is uniform with respect to  $\theta$  in  $0 \leq \theta \leq \pi$ . Hence it is continuous, and so equal to 0 at the end-points  $\theta = 0, \pi$ . Term by term differentiation yields

$$\frac{d\delta}{d\theta} = \Sigma \left[ \frac{nz}{n^2 - z} H_n \cos n\theta \right],$$

since (9) makes this series uniformly convergent in  $0 \leq \theta \leq \pi$ .  $d\delta/d\theta$  is

\* Cf. Southwell (*loc cit*), § 8.

† Whittaker and Watson, "Modern Analysis," § 3.64.

‡ Whittaker and Watson, "Modern Analysis," § 9.21

continuous in  $0 < \theta \leq \pi$  for the same reason. A second differentiation shows that

$$\frac{d^2\delta}{d\theta^2} = \Sigma \left[ \frac{-n^2 z}{n^2 - z} H_n \sin n\theta \right],$$

whenever this series is uniformly convergent, i.e.,

$$\begin{aligned} \frac{d^2\delta}{d\theta^2} &= -z \left( \Sigma [H_n \sin n\theta] + \Sigma \left[ \frac{z}{n^2 - z} H_n \sin n\theta \right] \right) \\ &= -z(H + \delta), \end{aligned}$$

whenever the series for  $H$  and  $\delta$  are uniformly convergent.

The series for  $\delta$  is uniformly convergent in  $0 < \theta < \pi$ , and the series in  $H$  is uniformly convergent in any interval  $(\alpha, \beta)$  where  $0 < \alpha < \beta < \pi$ \*. Hence

$$\frac{d^2\delta}{d\theta^2} + z\delta = -zH \quad (3) \text{ bis}$$

at all points of  $(0, \pi)$ , except possibly the end-points, and the doubt as to end-points may be removed, since by the continuity of  $d\delta/d\theta$  and  $\delta + H$

$$\begin{aligned} \left( \frac{d^2\delta}{d\theta^2} \right)_0 &= \lim_{\theta \rightarrow 0} \frac{1}{\theta} \lim_{\alpha \rightarrow 0} \int_{\alpha}^{\theta} \frac{d^2\delta}{d\theta^2} d\theta \quad (0 < \alpha < \theta) \\ &= \lim_{\theta \rightarrow 0} \frac{1}{\theta} \lim_{\alpha \rightarrow 0} \int_{\alpha}^{\theta} -z(H + \delta) d\theta \\ &= -z(H + \delta)_0. \end{aligned}$$

A similar argument holds at the end-point  $\pi$ .

We have proved that if  $H$  is continuous the series (6) is zero at  $\theta = 0, \pi$ , and satisfies the equation (3) at all points of  $(0, \pi)$ . The proof is easily extended to the case when  $H$  has a finite number of points of discontinuity such as would correspond to loading by couples proportional to  $P$  and applied at intermediate points of the beam.

#### *Evaluation of the Function $F_H(\theta, z)$*

2.2 The variation of  $F_H$  as  $z$  passes from 0 to 1 is a measure of the deviation from linearity of the exact theoretical  $\delta, v$  curve. Equation (7) indicates that such variations are likely to be small, and it could be used in any particular case to give numerical values, when, however, the loading can be simply expressed analytically, the simplest method is to solve equation (2), or its equivalent (3), in finite terms.

\* Whittaker and Watson, "Modern Analysis," § 9.24

We have from (3)

$$\delta = \frac{-z}{D^2 + z} H(\theta),$$

where  $D$  means  $d/d\theta$  as before. Also

$$\frac{1-z}{z} \delta = H(\theta) - F_H(\theta, z) \quad (7) \text{ bis}$$

Therefore

$$\begin{aligned} F_H &= \frac{1}{H} \frac{1-z}{z} \frac{-z}{D+z} H, \\ &= \frac{1}{H} \frac{-1+z}{D^2+z} H, \end{aligned} \quad (10)$$

where the operator is to be interpreted so that  $\delta$  is 0 at  $\theta = 0, \pi$ .

The following notation will be adopted for the forms which  $F_H$  takes in particular conditions of loading. The suffix to  $F$  will designate the source of "equivalent eccentricity." If the suffix designating a load stands alone the load is supposed to remain constant while  $P$  varies, and if the suffix is followed by  $/P$  the load is supposed to be kept proportional to  $P$ . This will be made clear by the following examples, which relate to the simple symmetrical cases, some of common occurrence, used later in this paper as illustrations. —

$F_A$  corresponds with equal end-eccentricities  $A$

$F_w$	„	constant uniformly distributed transverse load $w$ ,
$F_{w/P}$	„	transverse load $w$ , as above, but proportional to $P$ ,
$F_R$	„	two constant concentrated transverse loads $R$ , applied at two given points symmetrical about the mid length $c$ fig 3 (a),
$F_{R/P}$	„	transverse loads $R$ as above, but proportional to $P$ (this is the loading usually adopted in spar- tests),
$F_Q$	„	equal constant terminal couples $Q$ ,
$F_{Q/P}$	„	terminal couples $Q$ , as above, but proportional to $P$ . (In accordance with the footnote to § 2 I, $F_{Q/P} = F_A$ )

2.21 Numerical calculation is facilitated by taking advantage of a simple relation between some of the  $F$ -functions, which we now proceed to derive. Let  $W$  represent any form of loading by transverse forces or couples, and let

$M$  be the resulting bending moment and (as before)  $m$  the deflection resulting from  $M$  in the absence of other load. Concentrating now on any one pair of values of the loads  $P$  and  $W$ , we may calculate the total resulting deflection ( $\delta'$ , say), either on the assumption that  $P$  and  $W$  have been gradually applied simultaneously in constant proportion, or on the assumption that  $W$  was applied first, up to its full value, and then kept constant while  $P$  was applied. On the former assumption  $H$  has in equation (2) the form  $N(x)$ , which is now  $M/P$ —that is,  $M/2P_1$ , and  $\delta'$  is the same as  $\delta$ . Hence we have from (7)

$$\frac{1-z}{z} \delta' = HF_H = \frac{M}{2P_1} F_{W/P}(z),$$

so that

$$\delta' = \frac{1}{1-z} \frac{M}{2P_1} F_{W/P}(z) \quad (11)$$

Equation (11) holds for all values of  $M$  and  $z$  greater than 0. Letting  $z$  tend to zero with  $M$  unchanged, we deduce by continuity

$$m = \lim_{z \rightarrow 0} \delta' = \frac{M}{2P_1} F_{W/P}(0) \quad (12)$$

If now we take the latter assumption that  $W$  was applied first and then kept constant,  $\delta$  is  $\delta' - m$ , and  $H(x)$  is  $m(x)$  in equation (2) hence, by (7),

$$\frac{1-z}{z} (\delta' - m) = HF_H = m F_W(z),$$

and

$$\delta' = \frac{1}{1-z} m [1 + z \{F_W(z) - 1\}] \quad (13)$$

Comparing (13) with (11) and (12), we obtain the relation

$$F_{W/P}(z) = F_{W/P}(0) [1 + z \{F_W(z) - 1\}], \quad (14)$$

or, on re-arrangement and insertion of the argument 0 as a reminder that the result is true for deflection measured at any point of the beam,

$$F_W(z, \theta) = 1 + \frac{1}{z} \left\{ \frac{F_{W/P}(z, \theta)}{F_{W/P}(0, \theta)} - 1 \right\} \quad (15)$$

The identity (15) may also be obtained by algebraic manipulation of the symbolic definition (10)

It will be remembered that the form of an error function depends only on the distribution with respect to  $z$  (or  $\theta$ ) of the contribution made to  $H$  by the

relevant source of equivalent eccentricity. In the above work  $F_w$  corresponds with a contribution to  $H$  which is distributed like the *deflection* due to  $W$  alone, this deflection is typified by  $m(x)$  in equation (2).  $F_{w/r}$  corresponds with a contribution to  $H$  which distributed like the *bending moment* due to  $W$  alone, this bending moment is typified by  $N(x)$  in equation (2).

It follows that the identity (15) holds between the functions corresponding with any two sources of equivalent eccentricity, if the contribution to  $H$  of the first source is distributed with respect to  $x$  like the *deflection which would result from bending moments distributed like the contribution of the second*. Since the relation of deflection to bending moment is (for beams of uniform section) similar to the relation of bending moment to load distribution, the words in italics may be replaced by "*like the bending moment which would result from transverse loading distributed like the contribution of the second*". Provided that  $W$  does not include terminal couples, we deduce that (15) also holds when  $F_w$  is replaced by  $F_{w/r}$  and  $F_{w/r}$  is replaced by the  $F$ -function which corresponds with an initial deflection having the form of the curve of load distribution  $W$ . Accordingly, if the last  $F$ -function is known, all that is required to find  $F_w$  is two successive applications of (15).

The foregoing method is illustrated by a tabular example, Table I, in which  $W$  is taken to be a uniformly distributed load  $w$ . The connection with loading by terminal couples is also shown. Each of the diagrams in the third column shows the distribution along the beam's length of the contribution to  $H$  appropriate to the row in which it stands. The diagram in one row shows the form of deflection (or of bending moment) due to the form of bending moment (or of loading) shown in the diagram immediately below. Consequently an error function in one compartment is obtained by (15) from the error function in the compartment immediately below. We have  $F_n \equiv F_{Q/1}$ , as already noticed, because the initial form of the centre line when the loading is by eccentric end load is similar to the distribution of bending moment which results from terminal couples, and  $F_{w/r} \equiv F_Q$ , because the distribution of bending moment due to uniform transverse load is similar to the deflection due to terminal couples.

2.22 The relation (15) fails when  $z = 0$ , and at this point a limiting process is necessary. Denoting by  $F_{n+1}(z)$  the function derived from  $F_n(z)$  in the way that  $F_w$  is derived from  $F_{w/r}$ , we may rewrite (15), divided by  $(1-z)$ , as


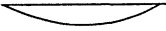

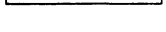
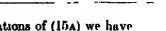
$$\frac{F_{n+1}(z)}{1-z} = \frac{1}{zF_n(0)} \left( \frac{F_n(z)}{1-z} - F_n(0) \right) \quad (15A)$$

Now assume that  $F_n(z)/(1-z)$  can be expanded in powers of  $z$ , that is, let

$$\frac{F_n(z)}{1-z} = c_0 + c_1z + c_2z^2 + c_3z^3 + \dots$$

The  $c$ 's are, of course, functions of  $\theta$

Table I

Source of equivalent eccentricity	Error function	Contribution to H	
Uniform load $w$ , $w/P = \text{constant}$	$F_w$		Deflection due to $w$
Uniform load $w$ $w/P = \text{constant}$	$F_{w/r}$		Bending moment due to $w$
End couples $Q$ , $Q/P = \text{constant}$	$F_Q$		Deflection due to $Q$
End-eccentricities	$F_A$		Graph of uniform load ing $w$
End couples $Q$ , $Q/P = \text{constant}$	$F_{Q/r}$		Initial position of centre line when end load is eccentric
			Bending moment due to $Q$

Then by repeated applications of (15A) we have

$$\frac{F_{n+1}(z)}{1-z} = \frac{1}{c_0} (c_1 + c_2z + c_3z^2 + \dots)$$

$$\frac{F_{n+2}(z)}{1-z} = \frac{1}{c_1} (c_2 + c_3z + c_4z^2 + \dots),$$

and so on, so that  $F_{n+1}(0) = c_1/c_0$ ,  $F_{n+2}(0) = c_2/c_1$ , etc. Thus we see that the values at  $z = 0$  of the functions derived from  $F_n$  are the ratios of consecutive pairs of coefficients in the expansion of  $F_n/(1-z)$  in powers of  $z$ .

2.23 Returning to our particular example, we can easily show, when  $H$  is made a constant in (3), that the form of  $F_H(\theta, z)$  as given by (7) or (10)—that is, of  $F_A$  (§ 2.2)—is given by

$$F_A(\xi, z) = (\cos \frac{\sqrt{z}\xi\pi}{2} \sec \frac{\sqrt{z}\pi}{2} - 1) \frac{1-z}{z}, \quad (16)$$

where  $\xi$ , which equals  $x/l$  or  $(\theta - \frac{1}{2}\pi)/(\frac{1}{2}\pi)$ , defines the point on the beam at which  $\delta$  is measured

By taking the limit  $z \rightarrow 1$ , we have

$$F_A(\xi, 1) = \frac{4}{\pi} \cos \xi \frac{\pi}{2}$$

Also, on division of  $F_A$  by  $1 - z$  and expansion in positive powers of  $z$ , the first coefficient and the ratios of following coefficients yield the values

$$F_A(\xi, 0) = \frac{\pi^2}{8} (1 - \xi^2)$$

$$F_{w,r}(\xi, 0) = \frac{\pi^2}{9} (1 - \frac{1}{2}\xi^2)$$

$$F_w(\xi, 0) = \frac{\pi^2}{10} \frac{du}{d\xi} \frac{(61 - 14\xi^2 + \xi^4)}{1 - \frac{1}{2}\xi^2},$$

and with the aid of these and equations (16) and (15) Table II is easily constructed. The percentage variation shown in the last column is

Table II

	$z$	0	0.2	0.4	0.5	0.6	0.8	1.0	% variation
$\xi = 0$	$F_A$	1.23370	1.24081	1.24820	1.25217	1.25616	1.26447	1.27324	+3.205
	$F_{w,r}$	1.02808	1.02880	1.02955	1.02994	1.03034	1.03118	1.03205	+0.386
	$F_w$	1.00341	1.00349	1.00358	1.00362	1.00367	1.00376	1.00386	+0.046
$\xi = \frac{1}{3}$	$F_A$	1.09662	1.09778	1.09897	1.09957	1.10017	1.10140	1.10266	+0.350
	$F_{w,r}$	1.00524	1.00529	1.00534	1.00557	1.00539	1.00545	1.00550	+0.026
	$F_w$	1.00025	1.00025	1.00026	1.00026	1.00026	1.00026	1.00026	+0.001

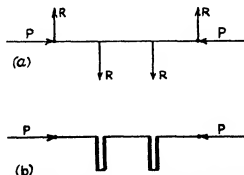


FIG. 3.—(a) Usual test loading. (b) Beam whose limiting form is the loading-curve of (a)

$100 \{F(1)/F(0) - 1\}$ . Values are shown for  $\xi = 1/3$  because at this point  $\sin 3\theta = 0$ , so that (if, as here, even harmonics in  $H$  are absent) only the fifth and higher odd harmonics in  $H$  affect the constancy of the  $F$ -function



2.24 As a further illustration of the method, some  $F$ -functions for the  $R$ -loading (§ 2.2 and fig. 3,  $a$ ) are tabulated below,  $\alpha$  is the distance from the ends of the points of application of  $R$ . These functions may be obtained by (15) from the limit ( $F_A$ , say) of the function corresponding with the beam shown in fig. 3,  $b$ , when the areas of the rectangular excursions remain constant as their breadth tends to zero. The vertical connecting parts, together with the right angle joints at their ends, are of course supposed rigid †. On taking the limit, we have

$$F_A \propto (1-z) \frac{2}{\sqrt{z}\alpha\pi} \sin \frac{\sqrt{z}\alpha\pi}{2} \cos \frac{\sqrt{z}\xi\pi}{2} \sec \frac{\sqrt{z}\pi}{2},$$

when  $\xi < 1 - \alpha$ ,

and hence, by expansion of  $F_A/(1-z)$

$$F_{R/P}(0) = \frac{\pi^2}{8} (1 - \xi^2 - \frac{1}{4}\alpha^2),$$

$$F_R(0) = \frac{5\pi^2}{48} \frac{(1 - \xi^2)(1 - \frac{1}{4}\xi^2 - \frac{1}{4}\alpha^2) + \frac{1}{4}\alpha^4}{1 - \xi^2 - \frac{1}{4}\alpha^2}$$

Table III

$z$		0	1	% variation
$\left. \begin{array}{l} \alpha = \frac{1}{2} \\ \xi = 0 \end{array} \right\}$	$F_{R/P}$	1.05003	1.05206	+0.193
	$F_R$	1.00186	1.00183	+0.007
$\left. \begin{array}{l} \alpha = \frac{1}{2} \\ \xi = \frac{1}{2} \end{array} \right\}$	$F_{R/P}$	0.91385	0.91189	-0.215
	$F_R$	0.99793	0.99785	-0.007
$\left. \begin{array}{l} \alpha = 1 \\ \xi = 0 \end{array} \right\}$	$F_{R/P}$	0.82247	0.81037	-1.447
	$F_R$	0.98696	0.98553	-0.145
$\left. \begin{array}{l} \alpha = 1 \\ \xi = \frac{1}{2} \end{array} \right\}$	$F_{R/P}$	1.05003	1.05206	+0.193
	$F_R$	1.00186	1.00183	+0.007

\* Since here  $\xi > 1 - \alpha$ , the functions have different analytical forms from those given above. It is easily verified that when  $F \equiv F_{R/P}$  or  $F_R$ ,  $F(\alpha, \xi, z) = F(1 - \xi, 1 - \alpha, z)$ . This is the extension of a well known reciprocal theorem to the case when  $P$  is present.

### Range of Application of the Linear $\nu - \delta$ Relation

2.3 From numerical results, of which the above are typical, it appears that in all ordinary cases in which either the loading is symmetrical, or in which  $\delta$

† It would perhaps be logically simpler to adopt a more cumbersome procedure, and obtain an algebraic expression for  $F_{R/P}$  from first principles by solving equation (3) with the appropriate (discontinuous) form for  $H$ . The result is, of course,

$$F_{R/P} = 1 + \frac{1}{z} \left\{ \frac{F_A(z)}{F_A(0)} - 1 \right\},$$

where  $F_A$  is as given above.

is measured so as to be unaffected by anti-symmetrical deflection, the variation in the  $F$ -functions is small, except possibly for  $F_A$ , it is less than could ordinarily be detected experimentally. Neglecting the variation, and noticing that the contributions to  $\delta$  of the various components of  $H$  are additive, we may summarize the results so far obtained as showing that an equation analogous to (1), viz ,

$$P_1 v - \delta = C \text{ (a constant),} \quad (17)$$

will hold very approximately in relation to a wide range of loadings. The constant  $C$  represents an "equivalent eccentricity" composed of—

- (i) the initial (unstressed) bowing at the point where the deflection is measured, multiplied by a factor which is nearly 1 if the bowing is nearly in the form  $\sin \theta$ ,
- (ii) the mean eccentricity of end-load, multiplied by a factor which is 1.25 (a mean value of  $F_A$  in Table II) for mid-length deflection,\*
- (iii) the deflection produced without end-load by any transverse loads or end couples which remain constant during test, multiplied by a factor nearly equal to 1,
- (iv) equivalent deflections corresponding with any transverse loads or end couples which are kept proportional to  $P$  during test, these deflections are independent of the flexural rigidity, being given by

$$\frac{\text{(Bending moment due to such transverse load or end couples at the point where the deflection is measured)}}{\text{(End load)}} \times \begin{cases} \text{a factor depending upon} \\ \text{the distribution of the} \\ \text{loads, and usually lying} \\ \text{between 0.8 and 1.25.} \end{cases}$$

Equation (12) shows that these are the deflections produced without end load by transverse loads or end couples proportional to the Euler critical end load,  $P_1$ .

In tests made by standard methods on aeroplane spars, the transverse loading is by intention entirely proportional to  $P$ , hence, apart from quantities of the nature of small corrections which can be estimated,  $C$  arises from (iv) above and depends on known quantities only. The experimental  $\delta, v$  points, therefore, may be expected to lie on a line which is not merely straight but also passes through a known point  $(-C, 0)$ . This fact adds to the already noticed

\* This is in effect a well-known result, given by Perry ('The Engineer,' vol 62, pp. 464 and 512 (1896)), where a value 5/6 (clearly a misprint for 6/5) is suggested as equivalent to our  $F_A$ .

advantages of the possibility of linear plotting, for example, an equivalent flexural rigidity corresponding to only one experimental point may be obtained from the slope of the line joining this point to the theoretical point  $(-C, 0)$

Equation (12) deserves attention, since,  $F_{w/r}$  being usually nearly unity, it shows that  $M/P_1$  is a fair approximation to the deflection due to the bending moment  $M$ . Equation (12) might also be derived as the limit of (10) as  $z \rightarrow 0$

#### *A Second Approximation to the $v - \delta$ Relation*

2.4. Before passing to experimental examples it may be of interest to examine further the nature of the approximation made in deriving equation (17)

2.41 We have, in effect, neglected all terms but the first in the numerator of the expression for  $F_n$  which is given by (7). If one more term is taken into account, say, the  $r$ th, then

$$F_n(z) = F_n(1) - \frac{1-z}{1-z/r^2} \{F_n(1) - F_n(0)\}$$

In the case of mid-length deflection the modifying term will ordinarily arise from the third harmonic, so that  $r^2 = 9$ . More generally, the approximate formula

$$F_n(z) = F_n(1) - \frac{1-z}{1-kz} \{F_n(1) - F_n(0)\}, \quad (18)$$

may be adopted, the constant  $k$  being adjustable. In (18)  $F_n(z)$  is supposed to be one of a series of functions connected as before by (15)

Equation (18) gives  $F_n$  truly at  $z = 0$ ,  $z = 1$ , and a value,  $k(z_1)$  say, may be assigned to  $k$  in order to ensure exact agreement at any chosen third point  $z = z_1$ . As  $z_1 \rightarrow 0$  or  $1$ ,  $k(z_1)$  tends to the value of  $k$  necessary to ensure agreement of the derivative  $F'_n$  at  $z = 0$  or  $1$  respectively. On taking limits and using (15),

$$k(0) = \frac{F_{n+1}(1) - F_{n+1}(0)}{F_{n+1}(1) - 1}, \quad k(1) = 1 - \frac{F_n(1) - F_n(0)}{F'_n(1)}$$

The most appropriate value of  $k$  will ordinarily lie between  $k(0)$  and  $k(1)$

The function  $F_n$ , with its comparatively large variation, affords a convenient numerical example. Values of  $k(z_1)$  are given in Table IV for the case  $\xi = 0$

Table IV.

$z$	0	0.2	0.4	0.5	0.6	0.8	1.0	% variation
$k(z_1)$	0.12373	0.12363	0.12334	0.12334	0.12313	0.12293	0.12273	-0.807

Thus  $k(z_1)$  is practically independent of  $z_1$ . If a mean value, 0.123234, be given to  $k$ , the formula (18) yields values of  $F_\lambda$  agreeing to at least six significant figures with the true values shown in Table II.

2.42 If (18) be assumed true, the  $\delta, v$  relation (17) is replaced by

$$P_1 v - \delta = C_1 - \frac{1-z}{1-kz} (C_1 - C_0),$$

where  $C_0$  and  $C_1$  are the values of the right-hand side at  $z = 0$  and  $1$  respectively. Remembering the definitions  $v = \delta/P$ ,  $z = P/P_1$ , we write  $z = \delta/P_1 v$  in the above, and thus obtain

$$P_1 v - \delta = C_1 - \frac{P_1 v - \delta}{P_1 v - k\delta} (C_1 - C_0),$$

which becomes, on reduction,

$$(P_1 v - \delta - C_1)(P_1 v - k\delta + C_1 - C_0) = -C_1(C_1 - C_0) \quad (19)$$

In the  $\delta, v$  diagram this represents a hyperbola passing through the origin. A typical form is sketched in fig. 4.

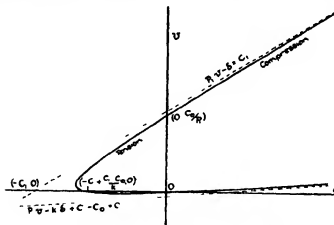


FIG. 4.—Hyperbolic approximation

### End Tension.

2.5 Hitherto  $P$  and  $z$ , and consequently  $\delta$ , have been assumed positive, attention being thus confined to the right-hand halves of figs. 1 and 4. The analysis is equally applicable, however, when  $P$  and  $z$  are negative, the end forces being then tensional. The solutions in finite terms now involve hyperbolic instead of circular functions thus  $\sec \sqrt{\epsilon} \pi/2$  in (16) is replaced by the

analytically identical form  $\text{sech } \sqrt{-z\pi/2}$  but the solution in series, as given by (7), is unchanged. The convergence of the numerator of the fraction in (7) being uniform in  $z$  for  $1 \geq z \geq -\infty$ ,\*  $F_A \rightarrow 1$  as  $z \rightarrow -\infty$ . Hence, when  $F_H(\theta, 0) \approx 1$ , the linear and hyperbolic approximations to the  $\delta, v$  relation may be expected to remain good for end tension. Actually the hyperbolic approximation (18), with the given value of  $k$  inserted, gives  $F_A$  with an error of less than 1% for end tensions less than  $15P_1$ , and less than 5% for all end tensions. Fig 5 reproduces the left-hand part of fig 4, and shows for this case (eccentric end load) both the true curve and the approximate hyperbola (19). The true value of  $dv/d\delta \rightarrow \infty$  as  $\delta \rightarrow -1$ .

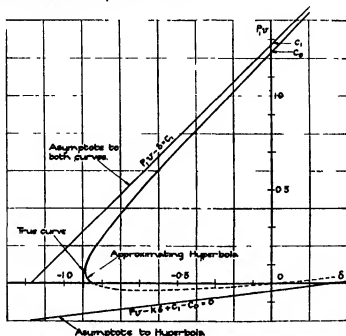


Fig 5—Hyperbolic approximation and ideal relation for end tension with unit end eccentricity

### Effects of Zero Errors

2.6 All of the preceding work depends on the assumption that  $\delta$  is the *additional* deflection—that is, the deflection which ensues when  $P$  and any loads proportional to  $P$  are applied. In actual tests, especially of “built up” spars,  $\delta$  may be subject to considerable zero error, of amount difficult to deter-

\* By Abel's test. Cf Bromwich, “Introduction to the Theory of Infinite Series,” § 44 (2), or Whittaker and Watson, “Modern Analysis,” § 3.35

mine directly, on account of minor settlements, etc., in the early stages of the loading process. It would not be fair to regard the effects of these as showing that the specimen differed materially in its behaviour from an ideal beam.

2.61 Let  $\delta$  be the true additional deflection in a test of an ideal beam, and let  $v = \delta/P$ . Let  $\delta'$  be the crude deflection (that is, the quantity measured instead of  $\delta$ ), and let  $v' = \delta'/P$ . Also let  $\delta = \delta' + \epsilon$ ,  $\epsilon$  being the constant correction which should be added to the measurements to give the true additional deflection.

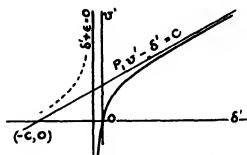


FIG. 6—Effect of zero error in  $\delta'$  ( $\epsilon$  positive).

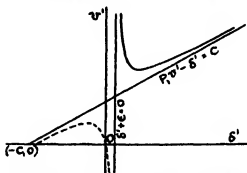


FIG. 7—Effect of zero error in  $\delta'$  ( $\epsilon$  negative).

Variation of the  $F$ -functions being neglected, we have

$$P_1 v - \delta = C,$$

that is

$$(\delta' + \epsilon) \left( P_1 \frac{v'}{\delta} - 1 \right) = C,$$

or

$$(P_1 v' - \delta' - C)(\delta' + \epsilon) = -C\epsilon \quad (20)$$

Hence the curve relating  $\delta'$  and  $v'$  is a hyperbola passing through the origin, of which the asymptotes are  $P_1 v' - \delta' = C$  and  $\delta' + \epsilon = 0$ . Figs 6 and 7 show the forms of this hyperbola for  $\epsilon$  positive and negative respectively.

2 62 In tests of actual material  $\epsilon$  will not be measured directly. If we assume that over a sufficient range the behaviour is like that of some ideal beam,  $\epsilon$  is easily found from three points on the corresponding range of the  $\delta', v'$  curve. Arithmetical methods may be used, but the geometrical method which follows has been found very satisfactory in practice.

2 621 Given  $O$  and three other points on the hyperbola (20), we wish to find the position of the vertical asymptote. Calling the three points 1, 2, 3, fig 8, we apply Pascal's hexagram theorem to  $O, 1, 2, 3$ , and the coincident points at infinity at which this asymptote meets the hyperbola, and obtain the

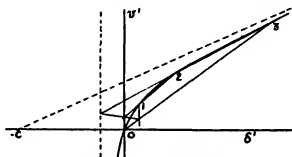


FIG 8.—Geometrical construction for zero error

following construction: the line joining the intersection of 2 1 and the ordinate through  $O$  with the intersection of 3  $O$  and the ordinate through 1 meets 2 3 at a point on the asymptote.

The other asymptote gives the required  $\delta, v$  relation, and may be obtained geometrically from the fact that any chord of a hyperbola makes equal intercepts between the curve and the asymptotes.

The corresponding arithmetical formula, and various "least-square" methods of solution, may be deduced without difficulty.

#### EXPERIMENTAL ILLUSTRATIONS

Some practical examples of the method are exhibited in figs 9 and 10.

The tests of fig 9 were undertaken with the special object of testing the theory. To avoid frictional errors at the end-pins the transverse loading was made self-balancing, both up and down forces being applied by means of a single dead weight connected to a system of levers, at each stage of transverse load the mid-length deflection was measured as the end-load was increased from zero up to a safe maximum. A test with eccentric end load only is included.

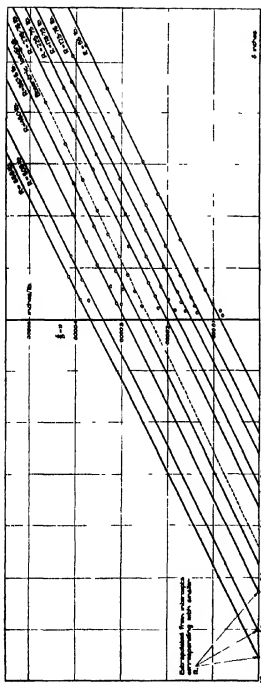


FIG. 9.—Solid rectangular steel bar  $\delta$  and  $\epsilon$  for (1) variable end load  $P$  with various constant values of transverse load  $R$ , as in fig. 3 (a) (full lines) (2) variable end load  $P$  applied eccentrically (broken line)

(1) for $R = 555 \text{ lb}$ $R = 505 \text{ lb}$ $R = 430 \text{ lb}$ $R = 367 \text{ lb}$ $R = 273 \text{ lb}$ $R = 223 \text{ lb}$ $R = 173 \text{ lb}$ $R = 123 \text{ lb}$ $R = 55 \text{ lb}$	$C = 0.6530$	extrapolated	$P_1 = 17,700$
	$C = 0.6025$		$P_1 = 17,665$
	$C = 0.5270$		$P_1 = 18,000$
	$C = 0.4788$		$P_1 = 18,170$
	$C = 0.3651$		$P_1 = 17,980$
	$C = 0.3306$		$P_1 = 18,020$
	$C = 0.2953$		$P_1 = 17,700$
	$C = 0.2332$		$P_1 = 18,000$
	$C = 0.1545$		$P_1 = 18,120$
			Weighted $\bar{P}_1 = 18,000$
(2) for eccentric loading $C = 0.442$			$P_1 = 18,480$



Fig 10 shows an airworthiness acceptance test of an aeroplane spar built up of steel strip, transverse and end loads being here proportional. Southwell's method of plotting reveals features of the test, such as the point at which the breakdown of elasticity begins, which it would be almost impossible to recognize by other methods. It is interesting to observe the similarity of these results to those obtained by Gough and Cox\* from tests on strip buckling under shear.

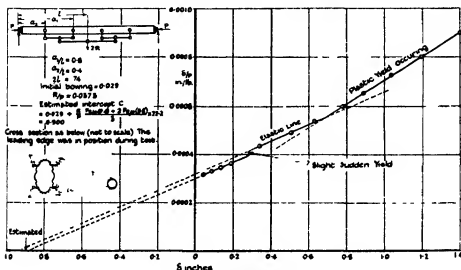


FIG 10—Aeroplane spar

#### Acknowledgments.

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#### Summary

The conventional theory of a beam subjected to combined transverse loads and axial end load is put into a form demonstrating an exact correspondence of transverse loads, either constant or proportional to end load, with initial (unstressed) deformation of the beam. A method of linear plotting suggested

\* *Loc cit*, fig 2

by Southwell for the analysis of experiments in elastic stability thus becomes available for the analysis of experiments in which "transverse" loads are operative and tend directly to increase the mode of deformation which becomes unstable. An examination in particular cases of the approximation implied by the linear plotting is facilitated by simple relations (derived from the above mentioned correspondence) between the functions expressing the error.

A very accurate second approximation is given, in which Southwell's straight line is replaced by a hyperbola.

Zero errors in the measurement of the deflection would destroy the collinearity of plotted observations, but methods are described for estimating and eliminating their effects

### *The Structure of Symm (1-3-5) Triphenylbenzene Part I*

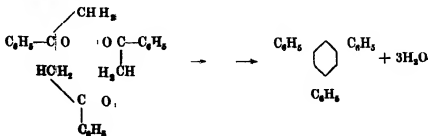
By BORIS ORELKIN and KATHLEEN LONSDALE

(Communicated by Sir William Bragg, O M, F R S—Received December 12, 1933)

*Note*—Boris Orelkin died on December 25, 1931. The X-ray investigation of symm. (1-3-5) triphenylbenzene was begun by him at the Davy Faraday Research Laboratory, where he worked as Research Fellow of the International Board of Education, but it was not completed at the time. The following paper contains his results and further preliminary data leading to an approximate structure. A full account of the detailed structure and of the magnetic properties will be published later.

#### *Chemical and Crystallographic Data*

The chemical structure of the symmetrical triphenylbenzene is determined by its preparation. It is obtained (cf the formation of mesitylene from acetone) by the action of dry HCl gas on acetophenone, HCl removing water according to the reaction



In the course of the reaction there are different intermediate products corresponding to the gradual condensation of acetophenone

The melting point of the crystals prepared in this way (by B O) was  $169^{\circ}$  to  $170^{\circ}$  C. A sample supplied later by Messrs Eastman Kodaks, and once recrystallized, melted at  $173^{\circ}$  C.

Very fine crystals of any weight from about 0.1 gm downwards can be obtained by crystallization from ether or ether + alcohol. These have exceptionally sharp bright faces, and the smaller ones sparkle like brilliants in a bright light. The larger crystals have a pronounced yellow tinge.

Symm triphenylbenzene has been investigated crystallographically by Knop, using crystals prepared by Arzruni. Their results are quoted by Groth† who gives diagrams of typical habits. The description and list of observed forms indicate that the crystals belong to the orthorhombic holohedral class, but, as will be shown later, they are, in fact, only hemihedral. The "imperfect cleavage" mentioned by Groth has not been observed on any of the crystals used in this investigation, a conchoidal fracture only being obtained. The habit and the supposed existence of this cleavage suggested to Fedorow that these crystals might be considered as "orthorhombic hexagonaloidal," and he gave them the habit symbol

$$\begin{array}{c} 6 \\ 37^{\circ} 44', \\ 0^{\circ} 25' \end{array}$$

where 6 denotes the "hexagonaloidal" character,  $37^{\circ} 44'$  the angle  $(1110 \ 1000)$   $[(10\bar{1}1) \ (0001)]$  in the usual notation and  $0^{\circ} 25'$  is the deviation from the hexagonal angle in the vertical zone  $[(010) \ (110)] - 60^{\circ}$ .

The habit is usually such that all the forms corresponding to the pseudo-hexagonal bipyramid and prism of the same order are represented, the orthorhombic nature of the crystals being indicated by the presence of other forms only occasionally developed. On the other hand, the development of the different planes in the pseudo-hexagonal forms is by no means even. In the crystallographic description the pseudo-hexagonal axis has been adopted as the *c* axis and the axial ratio is given as

$$a^* \ b^* \ c^* = 0.5662 \ 1 \ 0.7666,$$

the observed forms being  $\{110\} \ \{010\} \ \{011\}$  and less frequently  $\{112\} \ \{012\} \ \{100\} \ \{310\}$

† 'Chem. Kristallog.', vol. 5, p. 341 (1919).

Actually the so-called  $a^*$  axis of these crystals has now been found to be a polar axis, and, moreover, the  $c^*$  axis must be halved. Taking the polar axis as our new  $c$  axis (according to the usual crystallographic convention) the correct axial ratio becomes

$$a : b : c = \frac{c^*}{2} : b^* : a^* = 0.3833 : 1 : 0.5662$$

The planes (expressed in the correct notation, hereafter adopted throughout the paper) which have been observed during the course of the investigation now described are {011}, {010}, {120} and less frequently {001}, {111} and {013}.

{110} and {100} have also been observed, but at one end only. The {011}, {120} planes are always much better developed than the {010}, {111} planes, the

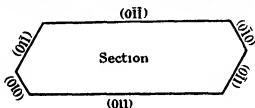


FIG. 1

last named being only occasionally developed. Usually the larger crystals are tabular on one pair of {011} faces.

The specific gravity, measured by Schroder,<sup>†</sup> is 1.205<sub>8</sub> (the exact temperature not stated).

On the basis of 4 molecules of  $C_8H_8$  ( $C_8H_8$ )<sub>4</sub>, molecular weight 306.14, in a unit cell of dimensions  $7.55 \times 19.76 \times 11.22 \text{ \AA}^3$ , the specific gravity is 1.205. These dimensions, obtained from rotation photographs taken at a plate distance of 10 cm, give an axial ratio

$$a : b : c = 0.382 : 1 : 0.568$$

#### *Space-group Determination*

Rotation and oscillation photographs about the three crystallographic axes, ionization spectrometer measurements and Weissenberg photographs of the three axial zones all agree in showing that the reflections from the following planes are absent —

(0*kl*) if ( $k + l$ ) is odd,

(*h*0*l*) if  $h$  is odd

<sup>†</sup> 'Ber. deutsch. chem. Ges.', vol. 14, p. 2516 (1881)

The absence of these reflections is characteristic of the space-groups —

$Q_h^{16}$  ( $Pnam$ ) in the holohedral class, and

$C_{2v}^9$  ( $Pna$ ) in the hemihedral class

In order to decide between these alternatives, the crystals were tested for the pyroelectric and piezo-electric effects† in the following way. A crystal measuring about 3 mm across was placed in a Pyrex test-tube and gripped between two insulated copper wires so that the latter made as good contact as possible with opposite (001) faces. The wires passed out through the cork of the test-tube, one being connected to earth, the other to the leaf of a sensitive Wilson tilted electroscope‡. When the surrounding test-tube was cooled by means of liquid air, the temperature of the crystal fell quickly and a small charge was developed at each end of the [001] axis. The leaf of the electroscope moved 10–12 divisions in repeated experiments. If both wires are earthed while the crystal is being cooled down, and then the same wire is connected to the electroscope, an opposite charge is indicated as the temperature rises. Reversal of the wires, of course, reverses the whole effect. Using a crystal of rock salt as a control, under similar conditions, a maximum deflection of 1 division was observed. Crystals of resorcinol and tartaric acid gave deflections of over 100 divisions, that is, greater than the whole length of the eyepiece scale. Since *symm.* triphenylbenzene shows this small, but positive, result for the piezo-electric (or pyro-electric) test, the space-group must be one in which there is *no centre of symmetry*.

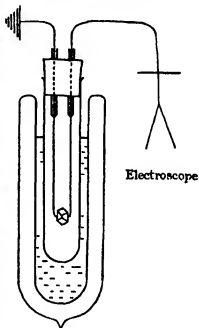


FIG. 2

† The strains set up by a change of temperature give a piezo-electric effect which is not distinguishable from a true pyro-electric effect. For references see 'Phil Mag.', vol. 6, p. 437 (1928).

‡ Sir William Bragg used an experiment similar to this in the course of a Royal Institution lecture, in order to demonstrate the polarity of resorcinol crystals.

The space-group therefore is  $Pna$ , in which there is a glide-plane of translation  $(b+c)/2$  parallel to (100), a glide-plane of translation  $a/2$  parallel to (010) and a two-fold screw axis parallel to [001]

The co-ordinates of equivalent points are

$$(x \ y \ z) \ (\bar{x} \ \bar{y} \ \frac{1}{2} + z) \ (\frac{1}{2} - x \ \frac{1}{2} + y \ \frac{1}{2} + z) \ (\frac{1}{2} + x \ \frac{1}{2} - y \ z)$$

the structure factor for any plane  $(hkl)$  being given by

$$|F(hkl)| = \sum f_o (A^2 + B^2)^{\frac{1}{2}}$$

(neglecting hydrogen atoms) where

$f_o$  = scattering power of carbon atom for plane  $(hkl)$

$$A = \sum \cos 2\pi (hx + ky + lz)$$

$$= 4 \cos 2\pi \left( hx - \frac{h+k+l}{4} \right) \cos 2\pi \left( ky + \frac{h+k}{4} \right) \cos 2\pi \left( lz + \frac{l}{4} \right),$$

$$B = \sum \sin 2\pi (hx + ky + lz)$$

$$= 4 \cos 2\pi \left( hx - \frac{h+k+l}{4} \right) \cos 2\pi \left( ky + \frac{h+k}{4} \right) \sin 2\pi \left( lz + \frac{l}{4} \right)$$

$$= 0 \text{ if } l = 0$$

The electron density at any point XYZ is given by

$$\rho(XYZ) = \frac{1}{V_c} \sum_{-\infty}^{\infty} \sum_{-\infty}^{\infty} \sum_{-\infty}^{\infty} |F(hkl)| \cos \left[ 2\pi \left( \frac{hX}{a} + \frac{kY}{b} + \frac{lZ}{c} \right) - \alpha(hkl) \right], \dagger$$

where the phase  $\alpha(hkl) = \tan^{-1} \frac{\sum B}{\sum A}$  (since there is only one kind of scattering material in the cell if hydrogens are neglected), and

$$\begin{aligned} F(000) &= \text{total number of electrons in the unit cell} \\ &= 96 \times 6 + 72 = 648 \end{aligned}$$

If  $l = 0$ ,  $\alpha(hkl) = 0$  ( $A$  positive) or  $\pi$  ( $A$  negative), that is to say, the projection of the structure on the plane (001) is centrosymmetrical. For this zone,  $(hkl)$ , the electron density is

$$\rho(XY) = \frac{1}{ab} \sum_{-\infty}^{\infty} \sum_{-\infty}^{\infty} F(hkl) \cos 2\pi \left( \frac{hX}{a} + \frac{kY}{b} \right)$$

$F(hkl)$  being positive or negative according to the sign of  $\sum A$ .

† Bragg, 'Z. Kristallog.', vol. 70, p. 476 (1929)

Since there are four molecules in the unit cell, each molecule is asymmetric. This was to have been expected since no symmetry has ever been found with certainty for the benzene nucleus other than a centre of symmetry† and, in one instance, a threefold axis‡. The formula of symm. triphenylbenzene precludes the possibility of a centre of symmetry, while the space-group forbids a threefold axis.

On the other hand the chemical formula strongly suggests a pseudo-trigonal axis in the molecule, in which case this would probably coincide, or make only a small angle with the pseudo-hexagonal crystal axis  $a$ . This arrangement can be tested by means of the optical and magnetic properties and by X-ray intensities.

### Optical and Magnetic Properties

The crystals show very strong negative birefringence, typical of the "layer lattice" structure type §.

The following data are given by Groth (*loc cit*) but are now referred to the correct axes.

The plane of the optic axes is (010),  $\alpha$  being the acute bisectrix. The refractive indices have the following values —

		$\alpha$	$\beta$	$\gamma$	
Li		1.5202	—	—	
Na		1.5241	1.8670	1.8725	
Tl		1.5291	1.8848	1.8897	
2V	9° 50'	Na	2E	17° 48'	Li
(calc.)	10° 17'	Tl	(obs.)	18° 25'	Na
				19° 27'	Tl

These values may be compared with the following —

	$\alpha$	$\beta$	$\gamma$
Naphthalene	1.442	1.775	1.932
Anthracene	1.43	1.79	2.04
Phenanthrene	1.335	1.538	1.625
Diphenyl	1.554	1.586	1.647
Hexamethylbenzene	1.503	1.747	1.801

† For references see 'Trans. Faraday Soc.', vol. 25, p. 352 (1929).

‡ Knaggs, 'Proc. Roy. Soc.', A, vol. 131, p. 312 (1931).

§ Wooster, 'Z. Kristallog.', vol. 80, p. 495 (1931).

In each of these structures the vibration direction of the fast ray (corresponding to the minimum refractive index) is perpendicular or most nearly perpendicular to the plane of the molecule, that of the slow ray (maximum refractive index) being parallel or nearly parallel to its longest dimension. With symm. triphenylbenzene it is clear that the plane of the molecule must be nearly perpendicular to the acute bisectrix  $a$  (that is, to the pseudo-hexagonal axis), while the arrangement of atoms in the (100) plane is such as to form an approximately uniaxial structure,  $\beta$  and  $\gamma$  being very nearly equal. The increased birefringence of triphenylbenzene relative to hexamethylbenzene points to an increased area of the molecule in its own plane.

Large optical birefringence in hydrocarbons is accompanied by strong diamagnetic anisotropy,<sup>†</sup> the (numerically) maximum diamagnetic susceptibility corresponding to the vibration direction of the fast ray. Experiment shows that for these crystals the susceptibility in the [100] direction is, in fact, very much larger than that along either of the other two axes, the susceptibility being a minimum (numerically) in the [001] direction. A more detailed account of the magnetic measurements will be published later.

### *X-ray Measurements*

A study of the measured intensities confirms the above molecular arrangement, in that the reflection from the (200) plane is very strong indeed, while the higher order reflections fall off steadily in intensity. The falling off is, however, more rapid than is consistent with an absolutely plane arrangement in the (200) planes. Moreover, if the atoms all lay in the (200) planes then  $x$  would always be 0 or  $\frac{1}{4}$ , giving

$$\begin{cases} A = 4 \cos 2\pi \frac{h+k+l}{4} \cos 2\pi \left( ky + \frac{h+k}{4} \right) \cos 2\pi \left( lz + \frac{l}{4} \right) \\ B = 4 \cos 2\pi \frac{h+k+l}{4} \cos 2\pi \left( ky + \frac{h+k}{4} \right) \sin 2\pi \left( lz + \frac{l}{4} \right), \end{cases}$$

if  $x = 0$ , or

$$\begin{cases} A = 4 \cos 2\pi \frac{k+l}{4} \cos 2\pi \left( ky + \frac{h+k}{4} \right) \cos 2\pi \left( lz + \frac{l}{4} \right) \\ B = 4 \cos 2\pi \frac{k+l}{4} \cos 2\pi \left( ky + \frac{h+k}{4} \right) \sin 2\pi \left( lz + \frac{l}{4} \right), \end{cases}$$

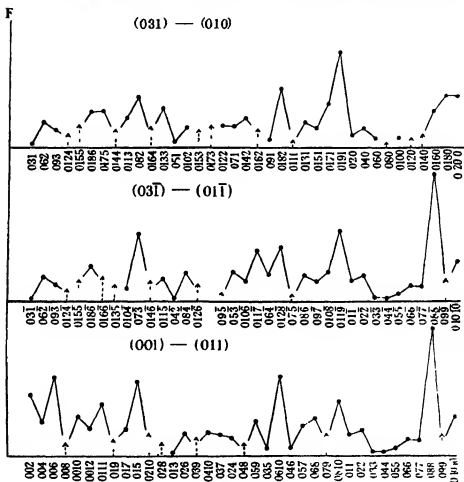
if  $x = \frac{1}{4}$

<sup>†</sup> Bhagavantam, 'Indian J. Phys.', vol 4, p 1 (1929)



In either case there would disappear certain sets of reflections (for example, from planes ( $hkl$ ) having  $h + k + l$  odd, in the first case) which have, in fact, been observed

Signs of pseudo-trigonal symmetry exist in the observed structure factors of the ( $0kl$ ) planes, fig 3 For example, the planes (015), (073), (082), which



hexagonal symmetry has almost entirely disappeared, so that a Laue photograph taken perpendicular to the  $[100]$  axis shows no trace of it. The slight tilt of the molecules to the  $(100)$  plane probably accounts partly for this effect, but it is also undoubtedly due to the introduction of the variable parameters consequent on reflection across the  $(100)$  and  $(010)$  planes. It is clear that even if the molecules were very nearly trigonal, and lay very nearly parallel to the  $(100)$  planes, suitable translations of the four molecules relative to each other would eliminate all pseudo-hexagonal symmetry from the structure as revealed by X-rays, though retaining it in the external form. The "centres" of the four molecules will lie at the points

$$(p \ q \ 0) \ (\bar{p} \ \bar{q} \ \frac{1}{2}) \ (\frac{1}{2} - p \ \frac{1}{2} + q \ \frac{1}{2}) \ (\frac{1}{2} + p \ \frac{1}{2} - q \ 0),$$

so that the  $(00l)$  planes are the only ones the structure factors of which are independent of  $p$  and  $q$ . Good agreement for all observed planes  $(00l)$ ,† up

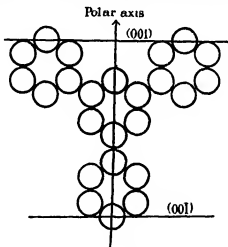


FIG. 4.

to  $l = 12$ , is obtained by placing a plane pseudo-trigonal molecule of 1-3-5 triphenylbenzene with one of its "arms" very nearly parallel to  $[001]$ , and in no other position. The polar axis of the crystal thus has a very real physical significance, owing to the fact that at one end of the crystal there are, in effect, twice as many carbon atoms in the surface as at the other end. Any redistribution of electron density due to temperature change or other cause of strain will have twice as much effect at one  $(001)$  face as at the other  $(00\bar{1})$  face.

† Mr B. W. Robinson very kindly made absolute measurements of the  $(002)$ ,  $(004)$ , and  $(006)$  reflections.

A detailed study of other hydrocarbons, notably of anthracene and naphthalene,† has shown that while the distance between the centres of carbon atoms in the same molecule is 1.54 Å or less, the corresponding distance between carbon atoms in different molecules is never less than 3.4 Å (as in graphite) and is usually about 3.7 Å for closest approach of neighbouring molecules ‡ Thus the van der Waals' forces between the molecules in the crystal may be represented by the enclosure of each carbon atom in a sphere of diameter 3.7 Å, within which no corresponding sphere from another molecule may be allowed to penetrate. This is a most useful rule, which can often be used to determine the most probable molecule arrangements in the crystal cell.

Taking the centre of the middle benzene nucleus as a representative point in the molecule, corresponding points in the four molecules are situated as follows —

Molecule 1	.	( <i>p</i> , <i>q</i> , 0)
Molecule 2 (reflection of molecule 1 in (100) <sub>1</sub> translated $\frac{1}{2}b + \frac{1}{2}c$ )		( $\frac{1}{2} - p$ , $\frac{1}{2} + q$ , $\frac{1}{2}$ )
Molecule 3 (reflection of molecule 1 in (010) <sub>1</sub> translated $\frac{1}{2}a$ )	.	( $\frac{1}{2} + p$ , $\frac{1}{2} - q$ , 0)
Molecule 4 (reflection of molecule 2 in (010) <sub>1</sub> translated $\frac{1}{2}a$ )		( $\bar{p}$ , $\bar{q}$ , $\frac{1}{2}$ ).

Consider the planes (*p*, 0, 0) and (1 + *p*, 0, 0), passing through eight molecules of type 1, which we know to be tilted a little out of the (100) plane, fig 5.

The molecule which occupies the space between four molecules of type 1 must be very near to the centre of gravity of the four, in order that the van der Waals' "spheres of force" shall not interpenetrate. This molecule can be either of type 2, or of type 4 (type 3 being in the plane ( $\frac{1}{2} + p$ , 0, 0)). If it is of type 2, then *p* and  $\frac{1}{2} - p$  must be nearly equal (their difference being not bigger than about  $\frac{1}{4}\epsilon$ ), while if it is of type 4, then *q* must be very nearly equal to  $\frac{1}{2}$  (say  $\frac{1}{2} \pm \frac{1}{4}\epsilon$ ). The latter arrangement would give a complete structure as shown in fig 6. This would not explain the big difference in the intensities of (002) and (031). It would, in fact, involve a much greater degree of pseudo-trigonal symmetry than is actually found, and so is out of the question.

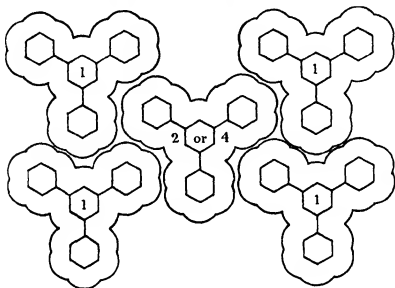
Placing a molecule of type 2 in this position, it is clear that the very small intensity of (031) is best explained by making *q* = 0,  $\frac{1}{2}$  or  $-\frac{1}{2}$  (nearly), so that if molecules 1 and 2 lie in the (031) planes, molecules 3 and 4 just interleave

† Robertson, 'Proc. Roy. Soc. A', vol. 140, p. 79 (1933), vol. 142, p. 674 (1933).

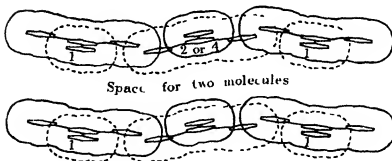
‡ Cf. also Müller, 'Proc. Roy. Soc. A', vol. 120, p. 455 (1928).

those planes, but all four molecules lie in successive (002) planes. These values of the parameters give arrangements as shown in fig 7

Fig 6 (a) shows that if  $q = 0$  the reflections from (011), (051), (015) planes should be very much weakened by comparison with those from (020), (042),



*a*



*b*

FIG 5

(082) which are inclined at  $60^\circ$  to them. This is not so, as fig 3 shows, and therefore we are left with  $p = \frac{1}{2} \pm (< 1\%)$  and  $q = \frac{1}{2}$  or  $-\frac{1}{2}$ , the differences in the corresponding structures being too small to be settled by any but the most detailed structure factor comparisons.

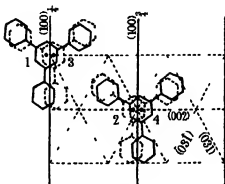


FIG 6

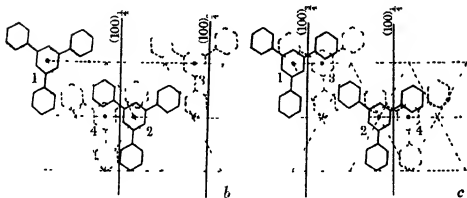
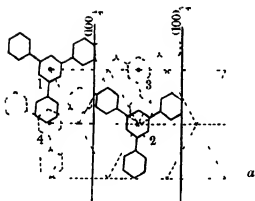


FIG 7

Discussion of further structural details, such as the exact molecular direction cosines, the distance between carbon atoms in a single nucleus and in different nuclei of the same molecule, will be left for the second part of this paper

#### *Acknowledgments*

One of the writers (B O) was indebted to the officials of the International Education Board for the grant of a science fellowship The other (K L.) sincerely thanks Sir Robert Mond, Sir William Bragg, and the Managers of the Royal Institution for financial assistance and laboratory facilities, and also for the continued interest they have shown in this work

#### *Summary*

1-3-5 triphenylbenzene crystallizes in the orthorhombic hemihedral class, space-group  $Pna$  ( $C_{2v}^8$ ) The unit cell contains four molecules and is of dimensions  $a = 7.55 \text{ \AA}$ ,  $b = 19.76 \text{ \AA}$ ,  $c = 11.22 \text{ \AA}$  The crystal habit is pseudo-hexagonal, with  $a$  as the pseudo-principal axis The optical and magnetic properties of the crystal and the X-ray measurements upon it show that the structure approximates to the "layer lattice" type Pseudo-trigonal molecules are inclined at a relatively small angle to the (100) planes, with one "biphenyl" arm nearly parallel to the [001] axis, so that there are, in effect, twice as many carbon atoms in the crystal surface at one end of the polar axis as at the other end The co-ordinates of representative points of the molecules are given to a first approximation

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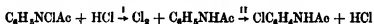
*The Effect of Solvent on Reaction Velocity V—The Interaction of N-Chloroacetanilide and Hydrobromic Acid in Dilute Aqueous Solution.*

By ISLWYN JONES and F G SOPER.

(Communicated by J L Simonsen, F R S —Received December 13, 1933)

The interaction of N-chloroacetanilide and hydrobromic acid to form *p*-bromoacetanilide and hydrochloric acid\* has, unlike the corresponding reaction with hydrochloric acid,† received comparatively little attention from a kinetic standpoint. It possesses, however, the advantage over the hydrochloric acid reaction, that its speed is much greater. It thus allows of the examination of mixtures containing only 0.001 M hydrobromic acid and of salt effects in the region of dilute concentration where the Debye-Hückel theory applies.

The mechanism of the interaction is analogous to that of the chloroamine-hydrochloric acid reaction, where the first stage



determines‡ the rate of transformation of the chloroamine in aqueous solution. Two corrections are necessary in this reaction if the true rate of interaction of chloroamine and acid is to be determined. The first correction is for the hydrolysis of the chloroamine under the influence of the acid, a rate mainly dependent on the H<sup>+</sup> ion concentration. Since the rate of the transformation depends on the product of the H<sup>+</sup> and Cl<sup>-</sup> ion concentrations, the correction for hydrolysis cannot be regarded as forming a constant fraction of the observed speed, particularly when the ratio of these ion concentrations is widely varied. The second correction is for the simultaneous formation of the chloroamines of the chloroanilides present. These chloroamines, which are formed pro-

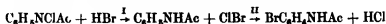
\* Acree and Johnson, 'Amer. Chem. J.', vol. 37, p. 410 (1907), Orton and Jones, 'Rep. British Ass.', p. 85 (1910), Richardson and Soper, 'J. Chem. Soc.', p. 1873 (1929).

† Orton and Jones (*loc. cit.*), Rivett, 'Z. phys. Chem.', vol. 82, p. 201, vol. 85, p. 113 (1913), Harned and Seltz, 'J. Amer. Chem. Soc.', vol. 44, p. 1476 (1922), Åkerlöf, 'Medd. Nobel Inst.', vol. 6, No. 2 (1925), Soper, 'J. Phys. Chem.', vol. 31, p. 1192 (1927), Soper and Pryde, 'J. Chem. Soc.', p. 2761 (1927), Fontein, 'Rec. trav. chim.', vol. 47, p. 635 (1928), Barnes and Porter, 'J. Amer. Chem. Soc.', vol. 52, p. 2973 (1930), Belton, 'J. Chem. Soc.', p. 116 (1930), Dawson and Millet, 'J. Chem. Soc.', p. 1920 (1933).

‡ Comparison of the observed rate of transformation with the value calculated for stage I from the equilibrium constant and the reverse velocity coefficient shows good agreement (Soper, *loc. cit.*)

gressively, appear in the chloroamine titre and cause the measured reaction rate to be slower by 4-10% than the actual rate of production of chlorine. This complication can be avoided by the presence of easily chlorinated substances, such as anisole, *p*-cresol,\* or acetanilide, which remove the chlorine as formed †

In the analogous hydrobromic acid reaction, the first stage



determines the rate in water. The correction for hydrolysis of the chloroamine is negligible in comparison with the measured rate of interaction, which is approximately 20,000 times faster than that of the hydrochloric acid reaction. The complication due to simultaneous formation of bromoamines by interaction of *p*-bromoacetanilide and ClBr also appears to be of much less magnitude and is completely eliminated by working in the presence of slight excess of acetanilide.

Before considering the salt effects observed in the present investigation at low ionic strengths, it is necessary to refer briefly to the conclusions reached from a study of the chloroamine-hydrochloric acid reaction. These conclusions, particularly those relating to the reactivity of the acid, depend largely on the fundamental reaction velocity equation adopted. Rivett (*loc cit*)

\* Belton (*loc cit*) regards the two corrections as eliminating each other and reports that he was unable to obtain consistent results using *p*-cresol. We have not experienced this difficulty when the *p*-cresol is quite colourless.

† Barnes and Porter (*loc cit*) have criticized this explanation of the reaction mechanism, particularly in regard to the effect of the addition of readily chlorinated substances on the rate of reaction. The experiments of Barnes and Porter were carried out in glacial acetic acid. In this medium the second stage determines the reaction rate (Orton and Jones, *loc cit*), the change of solvent having slowed up the interaction of chlorine and acetanilide and increased that of the chloroamine and hydrochloric acid. The state of affairs may be represented



where  $[\text{Cl}_2][\text{NHAc}]/[\text{HCl}][\text{NClAc}] = K$ . The reaction rate,  $v$ , is given by

$$v = k_2[\text{Cl}_2][\text{NHAc}] = k_2K[\text{HCl}][\text{NClAc}] = k_1[\text{NClAc}]$$

following a unimolecular law.

Addition of acetanilide increases both the concentration products  $[\text{HCl}][\text{NClAc}]$  and  $[\text{Cl}_2][\text{NHAc}]$  equally and increases the rate of disappearance of chlorine. The cause of the increase of the observed speed is different from that of the increased speed in water on addition of acetanilide. The presence of the more readily chlorinated  $\beta$ -acetanaphthalide in the glacial acetic acid medium, examined by Barnes and Porter, will obviously cause a more rapid disappearance of chlorine than the presence of a similar amount of acetanilide.



assigned separate reactivities to the un-ionized hydrogen chloride and to the  $H^+$  and  $Cl^-$  ions, using the classical equation in which velocity is strictly dependent on concentration. Harned and Seltz (*loc cit*) concluded that the speed was proportional to the chloroamine concentration and to the thermodynamic activity of the  $H^+$  and  $Cl^-$  ions. Soper and Pryde correcting for the two simultaneous reactions, previously referred to, found the speed proportional to the thermodynamic activity of both the chloroamine and hydrochloric acid molecules. Richardson and Soper suggested that this unusual concordance with the activity rate theory, might be associated with the constant surface tension and cohesion of the hydrochloric acid solutions examined. It was found, in the analogous reaction with hydrobromic acid that the activity rate theory held approximately for sucrose solutions, where there was little change in the cohesion of the solvent medium, but that divergencies were observed in opposite directions in aqueous alcohol and in salt solutions.

The agreement of results with the activity rate theory where the cohesion of the solvent medium remains constant is suggestive and may be due to the fact that under these conditions, the thermodynamic activities of the chloroamine and hydrochloric acid are proportional to the number of unsolvated or un-ionized molecules, respectively, able to pass a constant energy barrier at the interface at which the activity is determined, and thus measure the number of such molecules in unit free space. If the cohesion of the medium is not constant the energy barrier at the interface varies and the number of molecules escaping in unit time is not simply related to their concentration. The activity rate equation would appear to reduce to a true concentration reaction velocity equation when the medium is of constant cohesion and, further, this constancy results in uniformity of ease of formation or of resolution of the intermediate critical complex and results in the agreement observed.

Recently two papers have been published on the chloroamine-hydrochloric acid reaction by Belton and by Dawson and Millet, in which the results are regarded as supporting a theory of strict dependence of reaction speed on molecular concentration, over a range of considerable changes in the salt content of the reaction medium. According to Dawson and Millet, the reaction rate, being determined by the concentration of un-ionized hydrochloric acid, is proportional to the product of the concentrations of the  $H^+$  and  $Cl^-$  ions and their activity coefficients. The kinetic results are well accounted for if the activity coefficients are assigned values given by the equation,

$$\log f = -0.28\sqrt{\mu} + b\mu,$$

where  $b$  varies with the nature of the salt. These kinetically deduced values of the activity coefficients, however, differ considerably from those given by equilibrium measurements, and the discrepancy may be due partly to the non-correction of the observed speeds for simultaneous reactions and possibly also to a specific medium effect on the reaction process caused by the changes in the salt content of the solution.

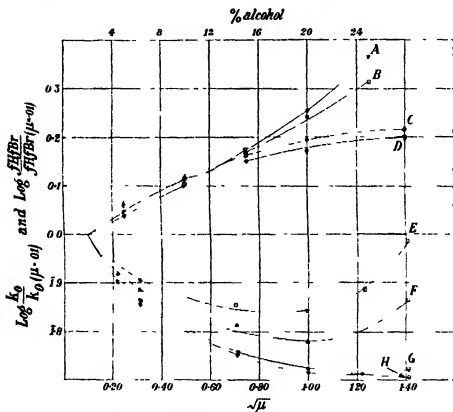


FIG. 1.—A,  $k_0$  in EtOH, B,  $k_0$  in MeOH, C,  $f_H f_{Br}$  in EtOH, D,  $f_H f_{Br}$  in MeOH, E,  $f_H f_{Br}$  in  $\text{NaNO}_3$ , F,  $f_H f_{Br}$  in  $\text{KNO}_3$ , G,  $k_0$  in  $\text{KNO}_3$ , H,  $k_0$  in  $\text{NaNO}_3$ .

Actual comparison of the velocity coefficients,  $k_0$ , of the hydrobromic acid reaction (where  $v = k_0 c_{H^+} c_{Br^-}$ ) with the measured thermodynamic activity coefficients of the  $H^+$  and  $Br^-$  ions in various aqueous media, was made by Richardson and Soper. These results are shown graphically in fig. 1, the relative change in the velocity coefficient caused by the various additions to the aqueous solution being compared with the relative change in the activity coefficients of the ions of the acid. These curves bring out the fact that in

sodium and potassium nitrate solutions the velocity coefficient,  $k_0$ , is depressed below the product of the activity coefficients whilst in alcoholic solution the reverse is true,  $k_0$  increasing more rapidly than the activity coefficient product. The specific rate of reaction of un-ionized hydrobromic acid with chloroamine is thus depressed in salt solution and increased in alcoholic solutions. These divergencies were regarded as specific solvent effects connected with the relative polar nature of the reagents and products and the cohesion of the solvent, which was regarded as assisting the intermediate critical complex to break up into those substances for which it has the greater affinity.

Comparison\* of the effect of ionic strength on the following types of ionic reactions



showed that increase of ionic strength, even in the range 0.001–0.03  $\mu$  increased the speed of reaction (1) above that of (2), as though the medium effect observed with neutral molecules were superimposed on the Bronsted primary salt effect. As high-valent ions were involved in the above comparison, with possible attendant anomalies, the investigation in dilute solution of the hydrobromic acid-chloroamine reaction, in which no ions have a higher charge than unity, was thought likely to be of interest.

Details of the method of following this reaction speed have previously been recorded. Sharp titration of the iodine liberated on addition of the mixture containing N-chloroacetanilide to aqueous potassium iodide was achieved, using N/500 thiosulphate and freshly prepared starch, by carrying out the operation in a current of nitrogen. The salts of the alkali metals and alkaline earths used as neutral salts in the investigation were B.D.H., A.R. chemicals and were dried to constant weight at 110°. The  $p_H$  of each solution was tested using brom-thymol blue and was shown to lie within the range 6.6 to 6.9. The nitric and perchloric acids were distilled from their constant boiling mixtures, diluted to the approximate strength required and estimated iodometrically. Conductivity water was used in all experiments. The velocity follows the equations for a bimolecular reaction, since in any one experiment, the  $H^+$  ion concentration is constant, i.e.,  $v = k_2 c_{NCl} c_{H^+} c_{Br^-} = k_2' c_{NCl} c_{Br^-}$ , where  $k_2 = k_2' c_{H^+}$ . The unusually good consistency of the bimolecular constants obtained may be illustrated by two typical experiments, recorded in Table I.

\* Soper and Williams, 'Proc. Roy. Soc., A', vol. 140, p. 59 (1933).

The collected results are given in Table II and for each series the slope of the  $\log k - \sqrt{\mu}$  curve has been calculated by the method of least squares. An estimate of the precision of the slope has been obtained by considering the

Table I

Initial concentrations  $[NCl] = [HBr] = 0.001 M$ ,  $[PhNHAc] = 0.01 M$ ,  $Ba(NO_3)_2 = 0.00500$ ,  $\mu = 0.01601$  thiosulphate = 1.044 N/500

Time (mins)	0	75	163	240	319	395	472	590
25 cc titre (cc thio)	21.28	17.51	14.55	12.64	11.12	9.84	9.00	7.98
$k_2$	—	3.14	3.16	3.16	3.16	3.15	3.13	3.16

Initial concentrations  $[NCl] = [HBr] = 0.001 M$ ,  $[PhNHAc] = 0.01 M$ ,  $[HClO_4] = 0.01501 M$ ,  $\mu = 0.01601$ , thiosulphate = 1.044 N/500

Time (mins)	0	5.00	10.00	15.00	20.00	25.91	33.00
25 cc titre	20.02	16.48	13.99	12.04	10.70	9.35	8.17
$k_2 \times 10^{-2}$	—	3.19	3.21	3.22	3.18	3.21	3.20

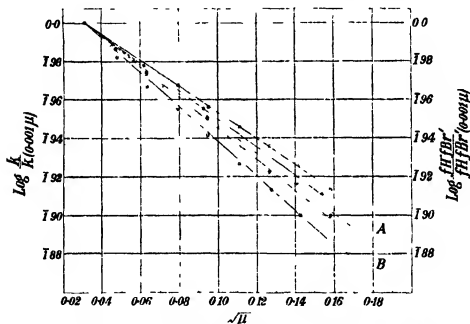


FIG. 2—O,  $HClO_4$ ,  $\square$ ,  $NaNO_3$ ,  $\bullet$ ,  $Ba(NO_3)_2$ ,  $\Delta$ ,  $HNO_3$ . A,  $f_H f_{Br} = f_H f_{Cl}$   
B,  $f_H f_{Cl} = f_H f_{Br}$  in KCl.

separate slopes of lines joining each point of the series to the calculated initial point of the series ( $\mu = 0.001$ ). The mean deviation of these slopes from the recorded value has been determined and the average error range of the slope of the series stated. In connection with each fresh salt series, a control

experiment on the reaction rate of chloroamine and hydrobromic acid, each at 0.001 M initial concentration, was carried out in the same thermostat and simultaneously with the fresh series of salt mixtures. The slopes of the curves, which appear to be linear up to  $\mu = 0.01$ , vary between  $-0.89$  and  $-0.64$  with an average precision of  $\pm 0.01$ . The results of four series are shown in fig. 2.

Table II

Salt added =  $\text{KNO}_3$  Initial concentration  $[\text{NCl}] = [\text{HBr}] = 0.001 \text{ M}$   
 Temperature =  $25.00 \pm 0.02^\circ \text{C}$

Cone of $\text{KNO}_3$	$\sqrt{\mu}$	$k_a$ range $\times 10^{-3}$	Mean $k_a$ $\times 10^{-3}$	No. of values of $k_a$	Log $k_a$
0.0000	0.03162	3.69-3.77	3.71	10	3.5694
0.00125	0.04793	3.54-3.60	3.56	10	3.5516
0.00299	0.06324	3.44-3.60	3.46	10	3.5389
0.005248	0.07905	3.37-3.40	3.38	12	3.5289
0.007999	0.09486	3.26-3.32	3.275	12	3.5152
0.011260	0.11070	3.19-3.18	3.15	12	3.4980
0.01501	0.12650	3.02-3.08	3.05	11	3.4843
0.01925	0.14230	2.92-2.99	2.95	12	3.4698
0.02400	0.15810	2.86-2.92	2.88	12	3.4594
0.03499	0.18972	2.69-2.74	2.715	12	3.4340
0.04900	0.22134	2.57-2.63	2.60	12	3.4150
0.06300	0.25296	2.50-2.54	2.52	12	3.4014
0.07996	0.28458	2.36-2.43	2.40	12	3.3793
0.09900	0.31620	2.33-2.38	2.35	12	3.3711
0.1200	0.34782	2.26-2.30	2.28	12	3.3570
0.1430	0.37944	2.21-2.27	2.24	12	3.3502

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.871 \pm 0.012$

## Addition of Magnesium Nitrate

Cone of $\text{MgSO}_4$	$\sqrt{\mu}$	$k$ range $\times 10^{-3}$	Mean $k_a$ $\times 10^{-3}$	No. of values of $k_a$	Log $k_a$
0.0000	0.03162	3.69-3.77	3.71	10	3.5694
0.0007475	0.06324	3.38-3.44	3.40	12	3.5318
0.00200	0.09486	3.10-3.14	3.115	12	3.4949
0.003782	0.12650	2.70-2.77	2.725	12	3.4354
0.0060000	0.15810	2.46-2.57	2.525	12	3.4023
0.008745	0.18972	2.26-2.32	2.285	12	3.3599
0.01300	0.22134	2.12-2.18	2.145	12	3.3324
0.01575	0.25296	1.87-1.92	1.89	12	3.2765
0.019999	0.28458	1.74-1.80	1.765	12	3.2467
0.02475	0.31620	1.59-1.65	1.628	12	3.2126
0.03000	0.34782	1.48-1.52	1.509	12	3.1786
0.03575	0.37944	1.39-1.44	1.417	12	3.1513

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-1.28$

Table II—(continued)  
Addition of Sodium Nitrate

Conc of $\text{NaNO}_3$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0 0000	0 03162	3 69-3 77	3 71	10	3 5694
0 00125	0 04743	3 53-3 58	3 575	10	3 5583
0 00299	0 06324	3 37-3 44	3 420	10	3 5354
0 005248	0 07905	3 36-3 39	3 38	10	3 5248
0 007999	0 09486	3 29-3 28	3 24	10	3 5105
0 011260	0 11070	3 10-3 15	3 13	10	3 4955
0 01501	0 12650	2 98-3 05	3 035	12	3 4821
0 01925	0 14230	2 91-2 97	2 9475	12	3 4694
0 02400	0 15810	2 86-2 90	2 88	12	3 4594

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.876 \pm 0.0082$ 

Addition of Lithium Nitrate

Conc of $\text{LiNO}_3$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0 0000	0 03162	3 68-3 75	3 71	10	3 5694
0 002493	0 05910	3 46-3 54	3 52	7	3 5465
0 005866	0 08870	3 31-3 38	3 34	7	3 5237
0 01300	0 1183	3 11-3 18	3 14	7	3 4999
0 02085	0 1473	2 93-2 99	2 95	7	3 4698

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.854 \pm 0.008$ Addition of Barium Nitrate Temperature =  $25.05 \pm 0.02^\circ \text{C}$ 

Conc of $\text{Ba}(\text{NO}_3)_2$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0 0000	0 03162	3 71-3 80	3 77	7	3 5763
0 000999	0 06324	3 53-3 58	3 55	7	3 5502
0 00266	0 09486	3 34-3 38	3 37	7	3 5276
0 00500	0 12650	3 13-3 16	3 15	7	3 4983
0 00800	0 15810	2 99-3 04	3 01	7	3 4786

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.7827 \pm 0.01$ Addition of Strontium Nitrate Temperature =  $25.05 \pm 0.02^\circ \text{C}$ 

Conc. of $\text{Sr}(\text{NO}_3)_2$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0 0000	0 03162	3 71-3 80	3 77	7	3 5763
0 000999	0 06324	3 53-3 59	3 54	7	3 5490
0 00266	0 09486	3 30-3 36	3 335	7	3 5230
0 00500	0 12650	3 09-3 15	3 13	7	3 4955
0 00800	0 15810	2 96-3 03	3 00	7	3 4711

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.7980 \pm 0.01$

Table II—(continued)

Addition of Perchloric Acid    Temperature =  $25.00 \pm 0.02^\circ \text{C}$ 

Conc of $\text{HClO}_4$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0.0000	0.03162	3.67–3.73	3.71	10	3.5694
0.00125	0.04743	3.50–3.57	3.55	7	3.5502
0.00299	0.06324	3.48–3.52	3.49	7	3.5428
0.006248	0.07905	3.39–3.47	3.44	7	3.5368
0.00799	0.09486	3.32–3.38	3.36	7	3.5283
0.011260	0.11070	3.26–3.29	3.27	7	3.5142
0.01501	0.12650	3.18–3.21	3.20	7	3.5051
0.01925	0.14230	3.08–3.16	3.13	7	3.4955
0.02400	0.15810	3.00–3.07	3.04	7	3.4829

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.0406 \pm 0.008$ 

Addition of Nitric Acid

Conc of $\text{HNO}_3$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0.0000	0.03162	3.67–3.73	3.71	7	3.5694
0.002829	0.06190	3.47–3.54	3.52	7	3.5468
0.007558	0.09251	3.30–3.36	3.34	7	3.5337
0.01420	0.12330	3.11–3.17	3.15	7	3.4983
0.02268	0.15390	3.00–3.04	3.02	7	3.4900

Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.741 \pm 0.008$ 

Addition of Sodium Chloride

Conc of $\text{NaCl}$	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0.0000	0.03162	3.67–3.75	3.73	7	3.5717
0.00299	0.06324	3.48–3.55	3.515	7	3.5459
0.00799	0.09486	3.33–3.39	3.27	7	3.5145
0.01501	0.12650	3.06–3.12	3.10	7	3.4914
0.02400	0.15810	2.84–2.89	2.87	7	3.4599

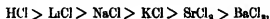
Slope of the  $\log k - \sqrt{\mu}$  curve =  $-0.8932 \pm 0.015$ Temperature =  $34.80 \pm 0.02^\circ \text{C}$ 

Conc	$\sqrt{\mu}$	$k_s$ range $\times 10^{-3}$	Mean $k_s$ $\times 10^{-3}$	No of values of $k_s$	Log $k_s$
0.000	0.03162	3.19–3.28	3.23	14	3.9154

No values for the activity coefficient of HBr in the presence of neutral salt at low ionic strength are recorded, but Livingston\* found that the activity coefficients of HCl and HBr are indistinguishable at concentrations below 0.1 M. According to the Bronsted theory of reaction velocity, the speed of the reaction will be given by  $v = C_{\text{H}} c_{\text{Br}} c_{\text{NCl}} f_{\text{H}} f_{\text{Br}} f_{\text{NCl}}/f_{\text{X}}$ , and in dilute solution by  $v = C_{\text{H}} c_{\text{Br}} c_{\text{NCl}} f_{\text{H}} f_{\text{Br}}$ . Comparison with the expression for the concentration velocity coefficient,  $k_0$ , shows that  $\log k_0$  should be given by

$$\log k_0 = \log C + \log f_{\text{H}} f_{\text{Br}},$$

and the curve of  $\log k_0 - \sqrt{\mu}$  should be exactly similar to one of  $\log f_{\text{H}} f_{\text{Br}} - \sqrt{\mu}$ . The values for the activity coefficient product are shown as a broken line curve in fig. 2, falling in the centre of the pencil of velocity curves. Data for the activity coefficients of hydrochloric acid† in the presence of neutral salts show that at constant ionic strength the salt effects are of similar magnitude to those observed kinetically, the divergencies which exist in the solutions of higher salt concentration tending to disappear in dilute solution. For purposes of comparison, the activity coefficient of hydrochloric acid in the presence of potassium chloride is indicated in fig. 2. A minor discrepancy exists in the order of the salt effects in the two cases. For the activity coefficients of hydrochloric acid at an ionic strength of unity, the order of  $f_{\text{H}} f_{\text{Cl}}$  is



whilst for  $k_0$  the order is

$\text{HClO}_4 > \text{HNO}_3 > \text{Ba}(\text{NO}_3)_2 > \text{Sr}(\text{NO}_3)_2 > \text{LiNO}_3 > \text{NaNO}_3, \text{KNO}_3 > \text{NaCl}$ , the relative positions of the alkali metal and alkaline earth salts being reversed. The great effect of magnesium sulphate on the reaction speed is largely due to removal of H-ion by the reaction  $\text{SO}_4^{2-} + \text{H}^+ = \text{HSO}_4^-$ .

The considerable variation in the slopes observed clearly demonstrates the specific nature of the salt effects and renders the comparison of the  $\log k - \sqrt{\mu}$  curves of ionic reactions of different types of uncertain value. In the present investigation with uni-valent ions, there is no evidence for a medium effect superimposed on the Bronsted primary salt effect in dilute solution. At higher salt concentrations the medium effect may possibly be expressed in the residual term  $f_{\text{NCl}}/f_{\text{X}}$  of the Bronsted equation, or, as previously suggested, may be due to an environmental influence assisting or preventing the break-up of the critical complex into the products.

\* 'J. Amer. Chem. Soc.', vol. 48, p. 45 (1926).

† Harned and Åkerlöf, 'Phys. Z.', vol. 27, p. 411 (1926).



In Table III the slopes of the  $\log k - \sqrt{\mu}$  curves are collected and compared with the relative cohesions of the various salt and acid solutions

Table III

Salt	Reciprocal of relative compressibility of 0.5 M solution	surface tension of solution at $\mu = 1.5$ $\Delta\gamma$	Slope of $\log k - \sqrt{\mu}$ curve
NaCl	1.178	2.45	-0.893 $\pm$ 0.015
NaNO <sub>3</sub>	1.139	1.76	-0.876 $\pm$ 0.006
KNO <sub>3</sub>	1.110	1.55	-0.871 $\pm$ 0.012
LiNO <sub>3</sub>	1.120	1.82	-0.854 $\pm$ 0.008
Nr(NO <sub>3</sub> ) <sub>3</sub>	—	1.27	-0.798 $\pm$ 0.010
Ba(NO <sub>3</sub> ) <sub>2</sub>	—	1.00	-0.783 $\pm$ 0.010
HNO <sub>3</sub>	1.020	-1.10	-0.741 $\pm$ 0.008
HBr	1.029	-0.69	—
HClO <sub>4</sub>	—	—	-0.641 $\pm$ 0.008

The surface tensions and the reciprocals of the relative compressibilities of the salt solutions indicate that the cohesions are in the order  $\text{NaCl} > \text{alkali nitrates} > \text{alkaline earth nitrates} > \text{acids}$ . This order is in agreement with the retarding effects of these solutions at constant ionic strength and in approximate agreement with the inverse order of the activity coefficients of hydrochloric acid in these solutions. At constant ionic strength, activity coefficients are regarded as affected by the average ion diameters and by the influence of the salts on the dielectric constant of the medium, but in view of the anomalous behaviour of the activity coefficients of the alkali hydroxides,\* where  $f_K > f_{Na} > f_{Li}$ , it is possible that other factors may sometimes dominate. If, for example, the equilibrium between co-valent hydrogen chloride and its ions varies in different solutions, this variation would be reflected in its activity coefficient. The effect of solvent on equilibrium is such that a change of solvent displaces the equilibrium,  $A \rightleftharpoons P$ , to form the substance, the solubility of which is the more increased by the change. By analogy with the behaviour of other co-valent gases,† increase of the internal pressure of an aqueous solution, which may be caused by the addition of salt, will decrease the solubility of co-valent hydrogen chloride and displace the equilibrium,  $\text{HCl} = \text{H}^+ + \text{Cl}^-$ , in the direction of the ions. In salt solutions at constant ionic strength,

\* Harned, 'J. Amer. Chem. Soc.', vol. 47, pp. 684, 689 (1925), Harned and Swindells, *ibid.*, vol. 48, p. 126 (1926).

† Euler, 'Z. phys. Chem.', vol. 31, p. 368 (1898), Hildebrand, "Solubility," N.Y. (1924).

increase of internal pressure would decrease the fugacity and the activity coefficient of the acid, as is observed. This effect would be additional to those from other causes. For the alkali hydroxides, an equilibrium of doublets with the free ions probably exists, and it is tentatively suggested that the anomalous order of activity coefficients for these substances, may be due to the opposite effect of internal pressure on the equilibrium of the ions, increase of internal pressure increasing the number of doublets and increasing the activity coefficient.

The critical increment of the reaction has been calculated from measurements at 25° and 35°, permitting the expression of  $k_0$ , referred to unit activity coefficient of the acid, by the equation

$$k_0 = 10^{14.39} e^{-14,580/RT}$$

The corresponding expression for the reaction of hydrochloric acid with the chloroamine, using the data of Harned and Seltz at 25° and 35° is

$$k_0 = 10^{14.00} e^{-21,370/RT},$$

the difference in the reactivity of the two acids being almost entirely due to the change in the critical increment from 14,580 to 21,370 cal. The products and reagents of contrasted nature in these reactions are hydrobromic acid and bromine chloride in the one reaction and hydrochloric acid and chlorine in the other. In both the contrast is similar, as also occurs in the chlorination of phenols, ethers and anilides,\* where chlorine and hydrochloric acid appear on opposite sides of the equation in all the reactions. This similar contrast in the nature of the products and reagents appears to confer constancy on the solvent effect in the series of reactions. In chlorinations, the changes in the reaction speeds are due, within experimental error, to changes in the energy of activation, as though the probability factor  $P$ , representing the influence of solvent on the ease of formation or of disruption of the critical complex into the products, is constant for the series. This factor, not unexpectedly, appears to be constant in the interactions of *N*-chloroacetanilide with hydrobromic and hydrochloric acids.

In conclusion, the authors wish to convey their thanks to the Chemical Society for a Research Grant, and to Professor J. L. Simonsen, F.R.S., for his continued interest in this work.

\* For references, *vide* Roberts and Soper, 'Proc. Roy. Soc.,' A, vol. 140, p. 71 (1933)

*Summary*

(1) The effect of neutral salt on the rate of interaction of hydrobromic acid and N-chloroacetanilide has been studied over the range of ionic strength, 0.001–0.025  $\mu$ , and the slopes of the  $\log k - \sqrt{\mu}$  curves, evaluated by the method of least squares, found to vary from  $-0.84$  to  $-0.89$  with the nature of the added electrolyte.

(2) The order of salt effects is approximately that of the activity coefficient of the acid and also that of the internal pressures of the solutions.

(3) The difference (20,000-fold) in the speeds of interaction of the chloro-amine with hydrochloric and hydrobromic acids is due almost entirely to the change in the energy of activation of the two reactions. This fact is regarded as supporting the interpretation of solvent effects previously advanced.

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*Optical Rotatory Power. I.—A Theoretical Calculation for a Molecule containing only Isotropic Refractive Centres*

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Revised February 23, 1934.)

*Introduction.*

Ever since the time of van't Hoff and Le Bel the number of investigations dependent on optical activity, or attempting to elucidate optical activity, has been very great, and it is remarkable that, even at the present time, there is no theoretical formula which gives the relation between the magnitude of the rotation and the chemical structure of the molecule concerned. The present communication supplies this want with regard to the molecule of the simplest asymmetric type—the molecule with four different groups attached to one central atom.

Various special hypotheses have been postulated to explain optical activity, but a few investigators have shown quite definitely that there is no necessity for any of these hypotheses. Born\* and Oseen† have shown independently

\* 'Phys. Z.', vol. 16, p. 261 (1915). 'Ann. Physik.', vol. 55, p. 177 (1918).

† 'Ann. Physik.', vol. 48, p. 1 (1915).

that, if the molecule has a dissymmetric structure, the ordinary refractive properties of the atoms will account for an optical rotation. Gray\* and de Mallemann† have attempted calculations of formulae for optical rotatory power on this basis. However, it has not been possible to condense the numerous algebraic terms which occur in these calculations into a compact form.

De Mallemann approached the nearest to the desired formula when he applied his calculation to a molecule of four atoms where one atom may be regarded as the origin of a set of rectangular axes and the other three atoms must lie on these axes. De Mallemann‡ showed that, if  $\text{CHClBrI}$  has a structure of this shape, the predicted rotation would be of the order of ordinary optical rotatory powers. However, there do not seem to be any compounds which are known to have the postulated structure, and the formula has no practical use.

By the following analysis it is possible to find a formula for the optical rotatory power of the general molecule of refractive centres, and this rotation has been expressed in a precise form by means of determinants. The expression is in a manageable form and simplifies by purely algebraic processes when it is applied to a particular type of molecule. It first became apparent to me that the ordinary properties of a dissymmetric molecule account for its optical activity, when I realized a complete analogy in a simple mechanical system. It is probably this analogy which has provided the clue for the solution of the complicated mathematical expressions which occur in the evaluation of the rotation caused by these refractive properties.

It is possible to construct a medium which shows a rotatory power for an ordinary transverse elastic wave. This is formed when we have an elastic solid with heavy masses embedded in it, as an example we may consider a gelatine gel with ball bearings as heavy particles. These balls are arranged in sets of four, all these sets are exactly alike, but the balls in each separate set may be different and may be at the corners of an irregular tetrahedron. These tetrahedral sets are distributed entirely at random, so that the gel as a whole is isotropic. There are two kinds of model, related to each other as an object to its mirror image, and hence a gel containing only one "enantiomorph" corresponds to an optically active liquid. Each ball vibrates under the influence of a passing wave and has an effect on neighbouring balls,  $e.g.$ , if a certain ball A vibrates up and down, then a smaller ball B a little farther

\* 'Phys. Rev.', vol. 7, p. 472 (1916).

† 'Rev. gén. sci.', vol. 38, p. 453 (1927).

‡ 'C. R. Acad. Sci. Paris', vol. 181, p. 298 (1925).

along the path of the wave, a little to one side, and a little above A, is dragged to and fro in a horizontal direction by the motion of A. The secondary wavelets caused by all these vibrators have to be added to the original wave and, if the total effect of these is not exactly in the same direction as the original wave, the plane of polarization is altered. For a medium containing a symmetrical model the resultant of these secondary wavelets would be in the same direction as the inducing wave, but with an unsymmetrical model this condition does not hold and the medium is "mechanically active."

This model bears the same relation to the natural optically active molecule as the Sellmeier vibrators in the early theory of optical dispersion do to the

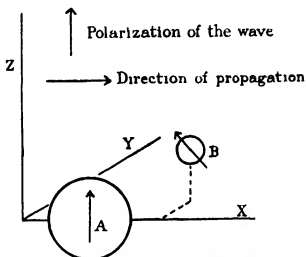


FIG. 1.—Direction of oscillation of B, a small ball embedded near a large ball A in an elastic medium

modern electron structure. Replace the ball bearings by atoms containing electrons free to move under the influence of an electric field, replace the elasticity of the gel by the equations of electricity, and the model becomes an asymmetric molecule under the influence of a light wave. Under the action of the electric field of a light wave each atom becomes an oscillating electric doublet. Inside the molecule the electric field of the wave is altered by the fields of the doublets themselves, and, making allowance for this, the exact polarizations of the molecule can be calculated. When the polarizations in a medium can be expressed in terms of the electrical fields, it is possible to find the velocity of a light wave in this medium by use of Maxwell's equations. Fresnel has shown that an active medium can be regarded as a medium in

which the velocities of circularly polarized waves of opposite sense are different, and the rotation can be calculated from this difference. Hence, having found the expression for the total polarization we can derive the formula for the rotation.

The method of evaluating the electric interactions of induced doublets in adjacent atoms has been employed successfully in other phenomena. W. I. Bragg,\* using the atom spacing found by X-rays, has calculated the various refractive indices of calcite and aragonite, and thus explained the strong double refraction of this type of crystal. Ramanathan† has deduced the anisotropy of various simple polyatomic molecules from the interaction of the atomic doublets, and has shown that the results agree with those obtained by the scattering of light.

To evaluate the electric interactions of the doublets it is necessary to know the size of these and the exact spacing of the atoms or groups in which they are induced. The size of an atomic doublet is obtained directly from the tables on atomic refractivity, but, while the spacing can be obtained directly for crystals, the shape of the active molecule has to be deduced from other data.

It is established now that the valency bonds in a molecule are easily flexible, and, in this calculation, it is assumed that the spacing in the molecule is very dependent on the close-packing of the atoms according to their size, the shape of the asymmetric molecule is taken as that determined by the close-packing of four spheres corresponding to the four different groups of the molecule.

The first stage of the calculation is carried out in this paper. A general expression is obtained for the rotatory power of any arrangement of polarizable centres in a molecule. Throughout the whole treatment Heaviside's rational units will be employed. In order to avoid repetition the theory of refraction according to Lorentz will be assumed.

#### *The Interactions of Induced Doublets in a Molecule and the Rotation of the Plane of Polarization*

The mechanical analogy suggests that the optical rotatory power may be due to the ordinary refractive properties of the atoms in the molecules, the interaction of the induced doublets can be calculated by the laws of electricity and the resultant electromagnetic properties of the whole molecule may be the

\* 'Proc. Roy. Soc. A,' vol. 105, p. 370 (1924).

† 'Proc. Roy. Soc. A,' vol. 107, p. 684 (1924).

origin of a rotatory power. It is necessary to calculate the electromagnetic properties of a fixed arrangement of polarizable particles in the molecule while neglecting all other chemical properties which do not interfere with the calculation. I have found it most convenient to define a hypothetical model and calculate exactly the relation between its dimensions and rotatory power, before examining how far its properties correspond to those of a chemical molecule. The generality of this treatment is so great that it shows that a model of conducting spheres joined together with insulating rods might twist the plane of polarization of a Hertzian wave, and that the amount of the twist could be calculated by the final formula.

We shall consider a medium with the following properties —

(1) The medium as a whole is perfectly isotropic, since it contains a large number of models distributed entirely at random.

(2) In accordance with the Lorentz treatment of refractivity, the field acting on any model is taken as  $E + aP$  where  $E$  is the electrostatic field of the electromagnetic wave,  $P$  the corresponding polarization and  $a$  is taken as  $\frac{1}{3}$ . This is tantamount to assuming that, owing to the random distribution, the medium external to the molecule is isotropic.

(3) The model consists of a rigid arrangement of particles which become polarized in an electric field. The particles are thus analogous to the refractive atoms in the ordinary molecule.

(4) The linear dimensions of the model are small compared with the wavelength of the electromagnetic wave.

(5) The polarization of each particle is proportional to, and in the same direction as the applied field when this field has a definite frequency.

(6) The particles are sufficiently symmetrical for the electric field of the induced doublet to be evaluated as if the doublet was concentrated at the centre of the particle.

(7) The changes in the doublets produced by the fields of local doublets are small compared with the magnitude of the doublet induced directly by the electric field of the electromagnetic wave.

When a single plane polarized wave acts on a model, an oscillating electric doublet is induced in each particle and, as one component of this may be out of phase with the inducing wave, we need two vectors to represent this. The electric field at a particle consists of the field of the light wave and the oscillating fields of the other doublets in the same model. To find the exact induced dipole in any one of a set of  $n$  particles it is necessary to solve  $6n$  equations in the components of these vectors. Later it is shown that the simplest active

molecule contains more than four atoms and so even for this molecule, there are twenty four equations for solution. A direct solution of these must be abandoned, but there is a method of evaluation which expresses the induced dipole as the sum of a series, every term of which is of the same kind and can be dealt with very simply, as shown in the following analysis.

The oscillating fields due to adjacent vibrators are small compared with the field due to the main light wave, and the magnitudes and directions of the vibrations can be obtained by successive approximations. The main field acting on a particle is that of the light wave, and the result is an induced dipole in phase with this. This initial dipole has an appreciable field at the positions of the adjacent atoms. On the initial dipole there are superimposed smaller dipoles due to the fields of initial dipoles of the adjacent atoms. This can be regarded as a second approximation and, passing to a third approximation we must consider the effects of these superimposed dipoles on the other vibrators, and so *ad infinitum*. The essential point is that the resultant dipole of any atom is obtained by the addition, vectorial with regard both to phase and direction, of a number of dipoles, each of which has been relayed by a definite path and has a definite source. Each of these is proportional to the dipole from which it originated and some constant depending on the course of the relays. Thus we can represent the vibration of A as made up of terms [A], [BA], [CA], [DA], [CBA], [ABA], [DCBA], etc., with these terms becoming less as their order increases.

If we resolve the initial dipole at A in three directions, every relay effect is included in the following type. If an induced dipole along the vector  $\alpha$  at A causes by means of the relay path [A B] a proportional induced dipole at B in the direction  $\beta$ . Letting AB be represented by  $\gamma$ , we have the whole system of interactions made up from terms of the type  $\alpha-\gamma-\beta$ . The effect of such a system on the difference in velocity of two oppositely circularly polarized waves can be found, and then this effect summed for all the systems in the molecule until these are of negligible size. It is most convenient to find the effect of this general relay system on the rotatory power before finding the exact value of any of the relay systems in the model.

When the model is situated in a variable electric field  $E$  with a value  $E_{A1}$  at A at a time  $t$ , the polarization along  $\alpha$  will be proportional to  $(\alpha \cdot E_{A1})$  and the relayed polarization to B can be written  $h\beta(\alpha \cdot E_{A1})$  where  $h$  is the constant of the relay system. Let  $\delta$  be the time lag during transmission. For the construction of the general electromagnetic equations of the medium it is necessary to express the polarization in terms of the electric field at the same point and



the same time. Since the model is small the relation between the field  $E_{A1}$  and the field  $E$  acting at  $B$  after  $\delta$  seconds is

$$E_{A1} = E - \frac{\partial E}{\partial x} \gamma_x - \frac{\partial E}{\partial y} \gamma_y - \frac{\partial E}{\partial z} \gamma_z - \delta E + \delta \frac{\partial E}{\partial x} \gamma_x + \delta \frac{\partial E}{\partial y} \gamma_y + \delta \frac{\partial E}{\partial z} \gamma_z \quad (1)$$

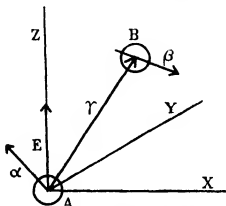


FIG. 2.—The relayed polarization

Substituting in the expression for relayed polarization we have the relation between the applied field and the polarization at the same point

The expression may be split into two similar parts dependent on  $E$  and  $-\delta E$  respectively. Since it will only be found necessary to consider relay terms of a low order, and the wave-length of the wave will thus be long compared with the relay path, the term  $-\delta E$  must be very small compared with  $E$ . The terms dependent on  $-\delta E$  will be neglected from this stage because if the analysis were completed it would be found that these have no appreciable influence on the rotatory power.

It is necessary to integrate the expression

$$h\beta [(\alpha \cdot E) - (\alpha \cdot \frac{\partial E}{\partial x}) \gamma_x - (\alpha \cdot \frac{\partial E}{\partial y}) \gamma_y - (\alpha \cdot \frac{\partial E}{\partial z}) \gamma_z] \quad (2)$$

where the vectors  $\alpha$ , etc., can have any direction so long as the relative configuration of these vectors remains the same. For the integration it is desirable to express the orientation in terms of independent variables, and this is performed by considering the position of the molecule as derived from the initial position  $\alpha_1 - \gamma_1 - \beta_1$  by the Eulerian rotations,  $\theta$  about the  $Y$  axis,  $\phi$  about the  $X$  axis, and  $\psi$  about the new position of  $\gamma$ . The rectangular axes  $X, Y,$

and  $Z$ , with the corresponding unit vectors  $i$ ,  $j$ , and  $k$  are taken in the usual positive sense so that  $i \wedge j = k$ . The position of any vector after performing the above rotations is obtained by operating with the tensor  $T$  whose coefficients are given in the table. The coefficients of the first row multiply the dyads  $ii$ ,  $ij$ , and  $ik$

$\cos \theta$	$\sin \psi \sin \theta$	$-\cos \psi \sin \theta$
$\sin \phi \sin \theta$	$\cos \psi \cos \phi - \sin \psi \sin \phi \cos \theta$	$\sin \psi \cos \phi + \cos \psi \sin \phi \cos \theta$
$\cos \phi \sin \theta$	$-\cos \psi \sin \phi - \sin \psi \cos \phi \cos \theta$	$-\sin \psi \sin \phi + \cos \psi \cos \phi \cos \theta$

(3)

If  $T\alpha_1$ , etc., are substituted in expression (2) we have the value of the polarization expressed in invariable vectors and functions of variable angles, and, since the number of molecules within the limits  $\theta$  to  $\theta + d\theta$ , etc., is

$$(N/8\pi^2) \sin \theta d\theta d\phi d\psi,$$

we can find the total value of the expression (2) for  $N$  molecules

For simplicity we shall carry out the integration of the various parts of (2) separately, and, taking the first term, we shall evaluate

$$(N/8\pi^2) \int_0^{2\pi} d\psi \int_0^{2\pi} d\phi \int_0^\pi 4T\beta_1(T\alpha_1 - E) \sin \theta d\theta \quad (4)$$

In the expansions of  $T\alpha_1$  and  $T\beta_1$  there are terms corresponding to the whole of the array of the tensor. Fortunately most of the coefficients become zero on integrating between the required limits, and we shall only construct the necessary vector terms after evaluating the coefficients. With regard to the integration  $\int_0^{2\pi} d\psi$  the expression is of the type

$$(A_1 + B_1 \sin \psi + C_1 \cos \psi) (A_2 + B_2 \sin \psi + C_2 \cos \psi),$$

and, after integration between the limits 0 and  $2\pi$ , the only terms which remain are due to  $A_1A_2$ ,  $B_1B_2 \sin^2 \psi$ , and  $C_1C_2 \cos^2 \psi$ . Only the corresponding terms remain in the integration  $\int_0^{2\pi} d\phi$ . In the third integration  $\int_0^\pi \sin \theta d\theta$  there is the extra factor  $\sin \theta$  and the terms which do not vanish are  $A_1A_2 \sin \theta$ ,  $(A_1B_2 + A_2B_1) \sin^2 \theta$ ,  $B_1B_2 \sin^3 \theta$ , and  $C_1C_2 \sin \theta \cos^2 \theta$ . By inspection of the product formed by squaring the array, it is seen that the only coefficients satisfying the above conditions are the squares of the original coefficients,

i.e.,  $a_{11}^2$ ,  $a_{12}^2$ , etc. Direct evaluation of these gives a total polarization of  $(N/3) hE(\alpha \cdot \beta)$ . The actual value of this is unnecessary in the calculation of optical rotation, but it is important that this portion of the polarization is in the same direction as  $E$ . This last result might have been considered obvious from considerations of symmetry.

Returning to the second term in the expression (2) we find that it is necessary to evaluate

$$-(N/8\pi^2) \int_0^{2\pi} d\psi \int_0^{2\pi} d\phi \int_0^\pi k T \beta_1 \left( T \alpha_1 \frac{\partial E}{\partial x} \right) \gamma_x \sin \theta d\theta \quad (5)$$

By the system of rotations adopted  $\gamma_x$  is equal to  $\gamma \cos \theta$ . With regard to the scalar coefficients in the expansion, this expression is the same as above, but with the extra factor  $\cos \theta$ . Proceeding as previously to find the characteristics of the terms which do not disappear on integration, we find that the conditions for  $\psi$  and  $\phi$  are as above, but only one term in  $\theta$  does not vanish i.e.,  $\cos^3 \theta \sin \theta$ . These conditions limit the coefficients to  $a_{22}^2 a_{22}$ ,  $a_{22}^2 a_{23}$ ,  $a_{22}^2 a_{33}$ , and  $a_{32}^2 a_{22}$ . The portions of these coefficients which do not vanish on integration are

$$a_{22}^2 a_{22} = -\cos^3 \psi \sin^3 \phi \cos \theta - \sin^3 \psi \cos^3 \phi \cos \theta$$

$$a_{22}^2 a_{33} = \cos^3 \psi \cos^3 \phi \cos \theta + \sin^3 \psi \sin^3 \phi \cos \theta,$$

giving

$$(N/8\pi^2) \int_0^{2\pi} \int_0^{2\pi} \int_0^\pi a_{22}^2 a_{22} \cos \theta \sin \theta d\theta d\phi d\psi = -N/6$$

$$(N/8\pi^2) \int_0^{2\pi} \int_0^{2\pi} \int_0^\pi a_{22}^2 a_{33} \cos \theta \sin \theta d\theta d\phi d\psi = N/6$$

The value of integral (5) is obtained by constructing the corresponding vectorial terms

$$\begin{aligned} & -\frac{Nh}{6} \left[ -j(k \cdot \beta_1)(j \cdot \alpha_1) \left( k \frac{\partial E}{\partial x} \right) \gamma + j(j \cdot \beta_1)(k \cdot \alpha_1) \left( k \frac{\partial E}{\partial x} \right) \gamma \right. \\ & \quad \left. - k(j \cdot \beta_1)(k \cdot \alpha_1) \left( j \frac{\partial E}{\partial x} \right) \gamma + k(k \cdot \beta_1)(j \cdot \alpha_1) \left( j \frac{\partial E}{\partial x} \right) \gamma \right] \\ & = +\frac{Nh}{6} \left[ k(j \cdot \frac{\partial E}{\partial x}) - j(k \cdot \frac{\partial E}{\partial x}) \right] [(j \cdot \beta_1)(k \cdot \alpha_1) - (j \cdot \alpha_1)(k \cdot \beta_1)] \gamma \quad (6) \end{aligned}$$

The last factor becomes

$$\gamma(k \wedge j \cdot \alpha_1 \wedge \beta_1) = -\gamma(j \cdot \alpha_1 \wedge \beta_1) = [\alpha_1 \gamma_1 \beta_1] = [\alpha \gamma \beta]$$

If the integrals corresponding to the other terms of expression (2) are calculated the only change is in the first bracket of the above expression, and these

factors together constitute curl  $\mathbf{E}$ . Thus the total polarisation due to  $N$  molecules containing the system of interactions  $\alpha - \gamma - \beta$  is

$$N\mathbf{E}(\alpha \beta) (\hbar/3) + N \text{curl } \mathbf{E} [\alpha\gamma\beta] (\hbar/6) \quad (7)$$

To obtain the total polarization of the medium this must be summed for all systems in all the molecules present, though it must be noted that many systems, while giving a value for the first term, will give a zero value for the triple scalar product in the second term.

Drude\* was aware that, when a term proportional to curl  $\mathbf{E}$  occurs in the expression for the average polarization of a molecule under the influence of an electromagnetic wave the medium containing this type of molecule will rotate the plane of polarization of the wave. He suggested that the electrons might be forced to vibrate in spiral paths, and after showing that this introduced a term of the type curl  $\mathbf{E}$ , he calculated the resulting rotatory power. However, there is not any other evidence of such a spiral system, and since the ordinary interactions of the doublets induced in the polarizable centres give a term of this type there is no necessity for the hypothesis of spiral movements.

For complete specification of the electrical properties, we require, in addition to the above relation, a knowledge of any magnetic induction due to the fields of the wave. In the ordinary refractivity treatment it is assumed that for the frequency of light waves the magnetic susceptibility is zero, and the molecule has no magnetic polarization. However, in this case there is a suggestion of a circular current in some of the polarizations of the molecule, this is especially apparent in the spiral motions which were introduced by Drude as a mathematical artifice, but which correspond mathematically to the present treatment. The possibility of a magnetic polarization has been frequently ignored in early examinations of this type, but recently it has been generally accepted. Gans† criticizing the analysis carried out by Born, has indicated the necessity for this, and the derivation of the value has been put in very clear form by Försterling‡. I had hoped to derive this magnetic moment directly from the spatial distribution of the oscillating polarizations in the molecule, but, though this method gives the same value, it is doubtful whether the method is absolutely rigorous. Hence, all justification for the use of this movement is based on the treatment as given by Försterling. When

\* "Theory of Optics," p. 401 (1902).

† "Ann. Physik.," vol. 79, p. 547 (1926).

‡ "Lehrbuch der Optik," Leipzig (1923), p. 133.

the energy of the wave is considered, it is shown that a term  $f \text{curl } \mathbf{E}$  in the electric polarization can only exist at the same time as a varying magnetic moment of value  $(f/c) \mathbf{E}$

Taking  $N$  as the Avogadro number and  $q$  as the concentration in gram-molecules per cubic centimetre we can write the electrostatic polarization  $\mathbf{P}$  and the magnetic polarization  $\mathbf{M}$  per cubic centimetre for any electromagnetic disturbance

$$\left. \begin{aligned} \mathbf{P} &= Ng \{ \mathbf{E} \Sigma h(\alpha \beta) / 3 + \text{curl } \mathbf{E} \Sigma h[\alpha \gamma \beta] / 6 \} \\ \mathbf{M} &= Ng \{ \mathbf{E} \Sigma h[\alpha \gamma \beta] / 6c \} \end{aligned} \right\} \quad (8)$$

The medium is perfectly isotropic and the only double refraction which can occur is a difference in the velocities of the two circularly polarized waves. We shall first find the refractive index of the medium for the circularly polarized wave

$$\mathbf{E}_1 = k\mathbf{E}_0 \cos\left(2\pi nt - \frac{2\pi x}{\lambda_1}\right) - j\mathbf{E}_0 \sin\left(2\pi nt - \frac{2\pi x}{\lambda_1}\right) \quad (9)$$

We have the polarizations expressed as dependent on the applied field, but according to the Lorentz treatment the applied field is equal to  $\mathbf{E}_1 + \alpha\mathbf{P}$ , where  $\mathbf{E}_1$  is the field of the wave. We can evaluate  $\mathbf{P}$  exactly for this particular wave, since  $\text{curl } \mathbf{E}_1 = 2\pi\mathbf{E}_1/\lambda_1$  and since  $\mathbf{P}$  is in phase with this we have  $\text{curl } \mathbf{E} = 2\pi(\mathbf{E}_1 + \alpha\mathbf{P})/\lambda_1$ . Substituting in equation (8) we obtain

$$\mathbf{E} = \mathbf{E}_1 / \{1 - aNg \Sigma(h(\alpha \beta) / 3 + 2\pi h[\alpha \gamma \beta] / 6\lambda_1)\} \quad (10)$$

To simplify write

$$\left. \begin{aligned} \epsilon &= Ng \Sigma h(\alpha \beta) / 3 \\ \rho &= Ng \Sigma h[\alpha \gamma \beta] / 6 \\ \phi &= 1 - aNg \Sigma(h(\alpha \beta) / 3 + 2\pi h[\alpha \gamma \beta] / 6\lambda_1) \end{aligned} \right\} \quad (11)$$

Substituting for  $\text{curl } \mathbf{E}_1$  by the second law of Maxwell  $\text{curl } \mathbf{E}_1 = -\mathbf{B}/c$ , equations (8) become

$$\left. \begin{aligned} \mathbf{P} &= (\epsilon\mathbf{E}_1 - \rho\mathbf{B}/c) / \phi \\ \mathbf{M} &= \rho\mathbf{E}_1 / c\phi \end{aligned} \right\} \quad (12)$$

Writing Maxwell's equations in the modified form for an isotropic medium we have the following equations to complete the solution, where  $\mathbf{D}$  is the electrostatic displacement,  $\mathbf{B}$  the magnetic induction and  $\mathbf{s}$  is unit vector in the direction of propagation

$$\left. \begin{aligned} \mathbf{D} &= \mathbf{E}_1 + \mathbf{P} & \mathbf{B} &= \mathbf{H} + \mathbf{M} \\ \mathbf{B} &= \mu\mathbf{s} \wedge \mathbf{E}_1 & \mathbf{D} &= -\mu\mathbf{s} \wedge \mathbf{H} \end{aligned} \right\} \quad (13)$$

$\mu$  is the refractive index and we can relate this to the electrical properties of the medium.

$$\left. \begin{aligned} \mu s_{\wedge} [\mu s_{\wedge} E_1] &= \mu s_{\wedge} B \\ &= \mu s_{\wedge} H + \mu s_{\wedge} M \\ &= D + (\rho/c\phi) \mu s_{\wedge} E_1 \\ &= E_1 (1 + \epsilon/\phi) + 2 (\rho/c\phi) \mu s_{\wedge} E_1 \end{aligned} \right\}, \quad (14)$$

but

$$\mu s_{\wedge} [\mu s_{\wedge} E_1] = \mu^2 s(s \cdot E_1) - \mu^2 E_1 = -\mu^2 E_1$$

Hence

$$E_1 (\mu^2 - 1 - \epsilon/\phi) = -2 (\rho/c\phi) \mu s_{\wedge} E_1 \quad (15)$$

Applying the relations solely to the circularly polarized wave represented in equation (9), whose field is  $E_1$  and whose particular refractive index is  $\mu_1$  we have,

$$s \wedge E_1 = -2\pi n E_1$$

$$(\mu_1^2 - 1 - \epsilon/\phi) = 4\pi\mu_1\rho/\phi\lambda,$$

$\lambda$  is the wave length of the light *in vacuo*

$$\left. \begin{aligned} (\mu_1^2 - 1) &= (\epsilon + 4\pi\mu_1\rho/\lambda) / [1 - a(\epsilon + 2\pi\mu_1\rho/\lambda)] \\ \frac{\mu_1^2 - 1}{a(\mu_1^2 - 1) + 1} &= \epsilon + \frac{2\pi\mu_1\rho}{\lambda} + \frac{2\pi\mu_1\rho}{\lambda} \frac{1}{a(\mu_1^2 - 1) + 1} \end{aligned} \right\} \quad (16)$$

This gives  $\mu_1$  the refractive index of the light corresponding to equation (9) but if the opposite kind of circularly polarized light is considered we obtain

$$\frac{\mu_2^2 - 1}{a(\mu_2^2 - 1) + 1} = \epsilon - \frac{2\pi\mu_2\rho}{\lambda} - \frac{2\pi\mu_2\rho}{\lambda} \frac{1}{a(\mu_2^2 - 1) + 1}$$

From these two equations we can obtain the value of  $(\mu_1^2 - \mu_2^2)$  by subtraction

$$\frac{\mu_1^2 - \mu_2^2}{[a(\mu_1^2 - 1) + 1][a(\mu_2^2 - 1) + 1]} = \frac{2\pi\rho}{\lambda} \left( \mu_1 + \mu_2 + \frac{(\mu_1 + \mu_2)[a(\mu_1\mu_2 - 1) + 1]}{[a(\mu_1^2 - 1) + 1][a(\mu_2^2 - 1) + 1]} \right)$$

The only term involving a difference of  $\mu_1$  and  $\mu_2$  is  $\mu_1^2 - \mu_2^2$  and, since the

difference in these refractive indices is very small, a mean value will be taken throughout the rest of the expression

$$\mu_1 - \mu_2 = \frac{2\pi Ng \sum h[\alpha\gamma\beta]}{6\lambda} [1 + a(\mu^2 - 1)][2 + a(\mu^2 - 1)] \quad (17)$$

The circularly polarized waves to which  $\mu_1$  and  $\mu_2$  apply respectively are given by the equations

$$\left. \begin{aligned} E_1 &= kE_0 \cos\left(2\pi nt - \frac{2\pi x}{\lambda_1}\right) - jE_0 \sin\left(2\pi nt - \frac{2\pi x}{\lambda_1}\right) \\ E_2 &= kE_0 \cos\left(2\pi nt - \frac{2\pi x}{\lambda_2}\right) + jE_0 \sin\left(2\pi nt - \frac{2\pi x}{\lambda_2}\right) \end{aligned} \right\} \quad (18)$$

Combining these two equations we obtain a plane polarized wave whose plane of polarization will rotate if  $\lambda_1$  and  $\lambda_2$  are unequal

$$\begin{aligned} E_1 + E_2 &= 2 \left[ kE_0 \cos \pi x \left( \frac{1}{\lambda_1} - \frac{1}{\lambda_2} \right) + jE_0 \sin \pi x \left( \frac{1}{\lambda_1} - \frac{1}{\lambda_2} \right) \right] \\ &\quad \times \left[ \cos \left( 2\pi nt - \pi x \left( \frac{1}{\lambda_1} + \frac{1}{\lambda_2} \right) \right) \right] \end{aligned} \quad (19)$$

Thus the vector determining the plane of polarization rotates on passing along the X axis. When  $x = 0$  the wave is polarized along the Z axis, but as  $x$  becomes larger the plane of polarization turns towards the Y axis, i.e., in a counter-clockwise direction when looking along the X axis, but in a clockwise direction when looking back along the X axis to receive the light. From the point of view of the observer the plane of polarization rotates  $(1/\lambda_1 - 1/\lambda_2)$  radians in a dextro or clockwise direction, and by equation (17)

$$\begin{aligned} \theta &= \pi l \left( \frac{1}{\lambda_1} - \frac{1}{\lambda_2} \right) = \frac{\pi l}{\lambda} (\mu_1 - \mu_2) \\ &= \frac{[a(\mu^2 - 1) + 1][a(\mu^2 - 1) + 2]\pi^2 Ng l \sum h[\alpha\gamma\beta]}{3\lambda^2} \end{aligned} \quad (20)$$

This is the rotation due to one compound of which there are  $Ng$  molecules per cubic centimetre when the light passes through  $l$  cm. of the medium of refractive index  $\mu$ .

This method of solution for the different refractive indices tends to be very formal, and it is advisable to consider the actual mechanism by which the difference arises. Considering the first wave (9) referred to above, we can resolve this into two plane waves travelling along the X axis. When the electrostatic field of the first is a maximum up the Z axis at the origin, the other

plane wave is a maximum at  $x = \lambda/4$  but acting along the Y axis. When part of the polarization of the molecule is dependent on curl E, not only does the Z wave cause a polarization up the Z axis, but it also causes a polarization a quarter of a period ahead and acting along the Y axis. If this extra polarization is positive, the polarizations in phase with each component of the circularly polarized wave are increased. In the circularly polarized wave of opposite sense the advance portion may be regarded as acting along  $-Y$ . Thus, while in one case the waves mutually reinforce each other by virtue of the curl E term, in the other they tend to annul each other. This causes a difference in effective dielectric constant and, neglecting magnetic effects, this gives an optical rotation. When the magnetic moment is taken into account the rotation is approximately doubled instead of the factor  $[\alpha(\mu^2 - 1) + 1]^2$ , which would be obtained if there were no magnetic effects, we have obtained the factor  $[\alpha(\mu^2 - 1) + 1][\alpha(\mu^2 - 1) + 2]$ .

The factor  $[\alpha(\mu^2 - 1) + 1][\alpha(\mu^2 - 1) + 2]$  has been retained until the present stage to illustrate the generality of the treatment and, though in future this will be written  $(\mu^2 + 2)(\mu^2 + 5)/9$  in accordance with the Lorentz theory, it would be quite possible to replace it by the appropriate factor in accordance with another theory of refractivity, such as that of Gladstone and Dale.

It may be noted that, even in entire absence of any chemical interaction, the rotation depends on the medium by means of the factor  $(\mu^2 + 2)(\mu^2 + 5)$ . In a mixture of two optically active substances the rotation will not obey a simple mixture law even if in every other respect the mixture is purely a physical one. The ordinary method of calculating the specific rotatory power, taking this as a property of the substance, tacitly assumes a simple mixture law and, though in the following analysis the value of the specific rotatory power is expressed, it must be recalled that this is only a constant for the compound under the special conditions of measurement. The specific rotation of a compound of which  $M$  g per cubic centimetre give a rotation of  $\theta$  radians in a column of length  $l$  cm is

$$[\alpha] = \frac{1800}{\pi g M l} \frac{\theta}{l},$$

where  $M$  is the molecular weight

The specific rotatory power of a compound in a medium of refractive index  $\mu$  is obtained from (20)

$$[\alpha] = \frac{(\mu^2 + 2)(\mu^2 + 5) 1800 \pi N}{27 M \lambda^2} \sum h [\alpha \gamma \beta], \quad (21)$$



where the summation  $\Sigma$  is carried out for every system  $\alpha - \gamma - \beta$  within the molecule

We have assumed that the exact magnitude of the doublets can be expressed in a converging series, and since these terms rapidly become more complex it will be very convenient if we can approximate at any stage by neglecting all higher terms. This is a definite approximation, but there is reasonable evidence in favour of this in the general optical properties of compounds—the fact that molecular refractivities can be calculated so accurately from atomic refractivities suggests that interactions between the doublets are small compared with direct polarizations, and the smallness of double refraction in crystals, compared with the mean refraction, indicates the relayed effects to be small compared with the primary polarizations. It will be shown presently that the optical activity is dependent on polarizations thrice relayed, and since the usual magnitude of optical rotations corresponds to a difference in the refractive indices of opposite circularly polarized waves of about  $10^{-6}$  it is apparent that the process of relaying decreases the magnitude of the polarizations considerably. Hence, when it is proved that a three relay system is responsible for an optical rotation, all higher relay terms will be neglected.

It is necessary to know the exact size of the electric doublet induced in any vibrator by a known electric field. In applying the formula to molecules these data can be obtained from the tables tabulated either by Bruhl or by Eisenlohr. Lorentz proved that, for a gram-molecule of a compound of refractive index  $\mu$  and density  $d$  the total polarization per unit applied field is  $3M(\mu^2 - 1)/(\mu^2 + 2)d$ , and it has been shown experimentally that each element makes a definite contribution, so that atomic polarizations can be found. The function tabulated, i.e.,  $A(\mu^2 - 1)/(\mu^2 + 2)d$ , where  $A$  is the atomic weight, represents one-third of the absolute atomic polarizability and is called the atomic refractivity. The moment induced in any chemical group is thus  $ER(3/N)$  where  $R$  is the sum of the atomic refractivities of the constituents of the group. The polarizability of the particle will be taken in this form for convenience in the application to chemical molecules.

In each particle of the model there is an oscillating dipole, and, since the wave-length of the electromagnetic wave is long compared with the distances at which the field acts on the other particles of the model, and since the dipole is considered as concentrated in a very small volume, the field is expressed by  $[1/4\pi r^2]\{3r(r \cdot D) - Dr^2\}$ . This is the simple electrostatic field at the

end of the vector  $r$  from a point dipole  $D$ . At  $A$  the initial dipole is  $E (3R_A/N)$ , and the field of this at  $B$  will cause a secondary dipole of value

$$\frac{9R_A R_B}{4\pi N^2 AB^3} [3AB (AB \cdot E) - EAB^2]$$

The first term of this represents a relayed effect which is proportional to the component of the field acting along  $AB$  at  $A$  and transmits a polarization along  $AB$  to  $B$  giving a polarization which also acts along  $AB$ . Thus using the general notation for a relayed effect we have  $\alpha = \gamma = \beta = AB$ , but since the effect on the optical activity is given by  $[\alpha\gamma\beta]$  it is apparent that the interactions responsible for a rotation must be much more complex than this.

The polarization  $[ABC]$  relayed to  $C$  is

$$\frac{27R_A R_B R_C}{(4\pi)^2 N^2 AB^2 BC^2} [9BC (BC \cdot AB) (AB \cdot E) - 3AB (AB \cdot E) BC \\ - 3BC (BC \cdot E) AB^2 + EBC^2 \cdot AB^2]$$

The polarization  $[ABCD]$  relayed to  $D$  is

$$\frac{81R_A R_B R_C R_D}{(4\pi)^2 N^4 AB^2 BC^2 CD^2} [27CD (CD \cdot BC) (BC \cdot AB) (AB \cdot E) \\ - 9BC (BC \cdot AB) (AB \cdot E) CD^2 \\ - 9CD (CD \cdot AB) (AB \cdot E) BC^2 + 3AB (AB \cdot E) CD^2 BC^2 \\ - 9CD (CD \cdot BC) (BC \cdot E) AB^2 + 3BC (BC \cdot E) CD^2 AB^2 \\ + 3CD (CD \cdot E) BC^2 AB^2 - ECD^2 \cdot BC^2 \cdot AB^2] \quad (22)$$

As might have been predicted from considerations of symmetry the relay effect involving the interactions of the four particles  $A$ ,  $B$ ,  $C$ , and  $D$  is the simplest effect to give a contribution to the rotatory power. Examining (22) we find that the 1st, 2nd, 3rd, and 5th terms give a definite value for  $[\alpha\gamma\beta]$ , *e.g.*, in the 1st term  $AB = \alpha$ ,  $CD = \beta$ ,  $AD = \gamma$ , and

$$h = \frac{81R_A R_B R_C R_D}{(4\pi)^2 N^4 AB^2 BC^2 CD^2} (CD \cdot BC) (BC \cdot AB)$$

A polarization  $h\beta (\alpha \cdot E)$  relayed along  $\gamma$  makes a contribution to the rotatory which is given by (21). The total rotatory power due to the relay path  $[ABCD]$  is

$$\frac{1800\pi (\mu^2 + 2) (\mu^2 + 5) 81R_A R_B R_C R_D}{27\lambda^2 MN^2 (4\pi)^2 AB^2 BC^2 CD^2} [27 [AB \cdot AD \cdot CD] (CD \cdot BC) (BC \cdot AB) \\ - 9 [AB \cdot AD \cdot BC] (BC \cdot AB) CD^2 - 9 [AB \cdot AD \cdot CD] (CD \cdot AB) BC^2 \\ - 9 [BC \cdot AD \cdot CD] (CD \cdot BC) AB^2] \quad (23)$$

All the triple scalar products are equal to  $\pm 6V$  where  $V$  is the volume of the tetrahedron  $ABCD$  and if we take  $[AB AD CD] = +6V$  then

$$[AB AD BC] = [BC AD CD] = -6V$$

The scalar products can easily be expressed in terms of the edges of the tetrahedron, *e.g.*,

$$(CD \cdot BC) = -CD \cdot BC \cos BCD = -(BC^2 + CD^2 - BD^2)/2$$

We can express the contribution of  $[ABCD]$  to the rotatory power completely in scalar quantities by these substitutions

$$\begin{aligned} & \frac{1800\pi (\mu^2 + 2) (\mu^2 + 5) 81R_A R_B R_C R_D}{27\lambda^2 M N^2 (4\pi)^3 AB^5 BC^5 CD^5} \\ & \times \frac{27V}{2} [3 (CD^2 + BC^2 - BD^2) (AB^2 + BC^2 - AC^2) \\ & - 2BC^2 (AD^2 + BC^2 - AC^2 - BD^2) - 2CD^2 (AB^2 + BC^2 - AC^2) \\ & - 2AB^2 (BC^2 + CD^2 - BD^2)] \quad (24) \end{aligned}$$

We shall only consider relayed effects of this type *i.e.*,  $[ABCD]$ , as the paths  $[ABCA]$ , etc., have no rotatory power and the higher terms such as  $[ABCDE]$  are assumed to be small compared with the terms of three relays. Hence for the total rotatory power of the model it is only necessary to consider all the combinations of four particles in the model and take all the relay paths within each set by considering all the permutations of these four.

We shall examine the rotatory power due to the interaction within the four particles  $A, B, C$ , and  $D$ . The rotatory contribution of any other path can be derived from the expression for  $[ABCD]$  by the obvious interchange of the letters, but the substitution must also be carried out in the triple scalar product represented by  $6V$  the value of this is perfectly constant, but the sign is changed. The triple scalar product for any path is found to have a value  $(-1)^n 6V$  where  $n$  is the number of inversions in the order of  $ABCD$  in the relay path.

The numerical value of  $V$  actually depends upon which "enantiomorph" is being considered and  $V$  will be considered as positive, thus making the whole analysis apply to a tetrahedral arrangement of the configuration shown in fig. 3. It can be noted that the configuration is defined by stating that when looking at the other particles from  $A$  these appear in a clockwise order in the order  $BCD$ .

The rule of signs for  $V$  is very suggestive of a simple method of expressing the sum of all these terms, since exactly the same rule of signs occurs in the expansion of a determinant. It appears as if there are twenty-four permutations to be considered, but owing to the symmetry of the expression, pairs of paths such as  $[ABCD]$  and  $[DCBA]$  give the same expression and there are only twelve different terms. Taking  $\Sigma \pm$  to mean the sum of all the terms

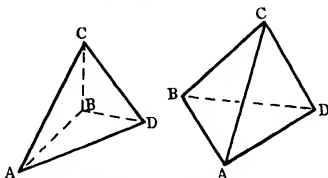


FIG. 3—Tetrahedra of the configuration which gives  $[AB AD CD]$  a positive value

with the signs chosen correctly we will omit the constants and consider only the following factor of the expression (24),

$$\begin{aligned} \Sigma \pm \frac{1}{AB^5 BC^5 CD^5} [BC^4 - 2AD^2 BC^2 + BC^2 (AB^2 + CD^2 - AC^2 - BD^2) \\ + 2AC^2 BD^2 - 2AB^2 CD^2 + (AB^2 CD^2 + AC^2 BD^2 - AC^2 CD^2 - AB^2 BD^2)] \\ = R^{-5} \Sigma \pm CA^5 AD^5 DB^5 (BC^4 - 2AD^2 BC^2 \quad \quad \quad), \quad (25) \end{aligned}$$

where

$$R^5 = AB^5 CD^5 AC^5 BD^5 AD^5 BC^5$$

Writing out the terms for some of the paths for  $\Sigma \pm CA^5 AD^5 DB^5 BC^4$ , the first term in the above expression,

$$\begin{aligned} [A B C D] + [D C B A] &= 2CA^5 DB^5 AD^5 BC^4 \\ [A C B D] + [D B C A] &= -2BA^5 DC^5 AD^5 BC^4 \\ [B A D C] + [C D A B] &= 2CA^5 DB^5 BC^5 AD^4 \\ [B D A C] + [C A D B] &= -2BA^5 DC^5 BC^5 AD^4 \end{aligned}$$

By addition we get  $2(AD^5 BC^4 + BC^5 AD^4)(AC^5 \cdot BD^5 - AB^5 \cdot CD^5)$

which is a minor of the following determinant, which is found to express all the terms of  $\Sigma \pm CA^5 AD^5 DB^5 BC^5$

$$\begin{vmatrix} 2 & 2 & 2 \\ (AB^5 CD^4 + AB^4 CD^5) & (AD^5 BC^4 + AD^4 BC^5) & (AC^5 BD^4 + AC^4 BD^5) \\ AB^5 CD^5 & AD^5 BC^5 & AC^5 BD^5 \end{vmatrix}$$

In future only the first columns of the determinants will be given as the succeeding columns can easily be constructed from these. Each of the other terms can be put into a corresponding determinant and the specific rotation due to the four particles A, B, C, and D is given by

$$[\alpha] = \frac{72900 (\mu^2 + 2) (\mu^2 + 5) R_A R_B R_C R_D I}{32 \pi^2 \lambda^2 MN^3}, \quad (26)$$

where

$$I = \frac{V}{R^3} \left\{ \begin{aligned} & \begin{vmatrix} 1 \\ AB^5 CD^4 + AB^4 CD^5 \\ AB^5 CD^5 \end{vmatrix} - 2 \begin{vmatrix} 1 \\ AB^7 CD^3 + AB^3 CD^7 \\ AB^5 CD^5 \end{vmatrix} \\ & + \begin{vmatrix} AB^2 + CD^2 \\ AB^5 CD^3 + AB^3 CD^5 \\ AB^5 CD^5 \end{vmatrix} - \begin{vmatrix} 1 \\ AB^5 CD^3 + AB^3 CD^5 \\ AB^5 CD^5 (AB^2 + CD^2) \end{vmatrix} \\ & + 2 \begin{vmatrix} 1 \\ AB^5 + CD^5 \\ AB^7 CD^7 \end{vmatrix} - 2 \begin{vmatrix} AB^2 CD^2 \\ AB^5 + CD^5 \\ AB^5 CD^5 \end{vmatrix} \\ & + \begin{vmatrix} 1 \\ AB^5 (AD^2 BC^2 + AC^2 BD^2 - AC^2 CB^2 - AD^2 DB^2) \\ + CD^5 (AD^2 BC^2 + AC^2 BD^2 - AC^2 AD^2 - CB^2 BD^2) \\ AB^5 CD^5 \end{vmatrix} \end{aligned} \right\} \quad (27)$$

The interactions of the four particles A, B, C, and D cause a rotatory power of  $[\alpha]$  and, if there are more than four particles in the model, the total rotatory power is obtained by summing all the contributions from all the combinations of four particles in a single model.

If there is a plane of symmetry in the tetrahedron ABCD, or for more than four particles if the model is symmetrical as a whole, it is apparent that the

rotatory power is zero. Regarding a molecule as an atomic model we see that the same conditions seem to apply to natural rotatory power, active molecules are always dissymmetric from a chemical point of view, and it is obvious that the difference in the sizes of the various atoms will make the spatial arrangement dissymmetric.

In a liquid we have an almost random distribution of molecules, each of which consists of a definite structure of atoms. The ordinary refractivity data show the various parts of the molecule to be polarized by the electric field of a light wave, and, if we use, as an illuminant, light of a wave-length considerably longer than X rays, whose wave-length is comparable with atomic dimensions, we have a molecular system fulfilling the properties of the hypothetical model used above. The conditions 5-7 were approximations made for the sake of simplicity in the mathematical treatment, but these will hold fairly accurately for a molecular system. Formula (26) gives the rotatory power of a chemical compound in terms of the atomic refractivities and the internal dimensions of the molecule.

A definite approximation has been made when the particles are considered as isotropic, and it is of special interest to examine this more closely since Kuhn\* attributes the rotatory power to the anisotropy of radicals in the molecule. In the present calculation it is not at all essential for the particles to be isotropic in order that the molecule be optically active, but our knowledge of anisotropy is very vague, and so this is a very convenient approximation. If we are to connect optical rotatory power with known data, we shall have to neglect this variation in refractivity with direction. In many simple molecules which are optically active there is no evidence of anisotropy, and since the anisotropy is not necessary for the existence of optical activity it can be classed as a secondary effect. Hence it is probable that even in more complex compounds the rotatory power will be given fairly accurately by considering only the average refractive properties.

By formula (26), the rotatory power of any molecule could be calculated. In a straightforward application of the formula to a given compound, each atom could be taken as a refractive centre. The distances AB, etc., would be the distances between the centres of pairs of atoms. Formula (26) gives the contribution of a combination of four atoms to the total rotatory power, and all the contributions (i.e., from every possible combination of four atoms) would have to be summed. Unfortunately the complexity of the formula makes such a procedure impossible, even if the calculation were practicable,

\* 'Trans. Faraday Soc.,' vol. 22, p. 457 (1926)

there is no direct way of measuring all the interatomic distances. The formula is only fit for a practical use if it can be applied to a particular type of atomic structure so that (a) it becomes much simpler in form, and (b) the distances AB can be easily expressed in terms of some atomic property which is already known.

The paper following this contains the account of the application of the formula to the simplest type of optically active molecule.

At the present stage we have established that the magnitude of the optical rotation as well as its sense is completely determined by the arrangement of refractive centres in the molecule. The optical rotatory power is expressed in a precise formula which must be applied to the various structural types of optically active compounds in order to obtain for them a practical theoretical formula.

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### *Optical Rotatory Power. II—The Calculation of the Rotatory Power of a Molecule containing Four Refractive Radicals at the Corners of an Irregular Tetrahedron*

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It has been shown in the previous paper that, where there are four refractive centres arranged dissymmetrically in a molecule, there is a definite rotatory power depending on the dimensions of the molecule. In the simple active molecule of four different radicals we seem to have been provided directly with four different refractive centres. It is, however, necessary to examine if a chemical radical fulfils all the properties which were assigned to the hypothetical particle in the general calculation, but, if this is true, we can substitute dimensions of such a molecule in the general formula to derive a simpler formula for the rotation.

If the radical is to have the same properties as the polarizable particle, it will have a definite refractivity for a fixed wave-length and its electric field in the polarized state must approximate to that of a dipole of equal value situated at its centre. A chemical radical has a definite refractivity for a given frequency, but it is only if the radical is small and fairly compact that its

field in the polarized state may be taken as that of a dipole concentrated at a single point. Consequently this paper will be limited to the calculation for molecules containing radicals of only a few atoms in length. The refractive properties will be regarded as concentrated at the geometric centre of these small radicals.

When different radicals are attached to a common atom, the Thorpe-Ingold theory postulates that the positions of the radicals are largely controlled by their respective volumes. This conception of atomic close-packing within the molecule has been found necessary to explain the physico-chemical properties of many compounds, and more recently it has been directly confirmed by the X-ray diffraction and electron diffraction experiments of Debye and Wierl\* respectively and their collaborators, these investigators have measured the dimensions of the molecules  $\text{CCl}_4$ ,  $\text{CHCl}_3$ , and  $\text{CH}_2\text{Cl}_2$  in the vapour state, and have shown the configuration to be determined by the sizes of the Cl and H atoms. It appears that radicals linked to the same central atom are drawn into contact and that the valencies have little or no directing effect. This is very convenient for the calculation of the shape of the tetrahedron of centres when four different radicals are present since the radicals are small we can regard each as having a sphere of repulsion and the distance between any two centres will be the sum of the radii, i.e.,  $AB = (a + b)$ , etc.

We shall proceed to calculate the rotatory power of four refractive centres of value  $R_A$ ,  $R_B$ ,  $R_C$ , and  $R_D$  situated at the apices of a tetrahedron whose edges have the lengths  $(a + b)$ ,  $(b + c)$ , etc. This rotatory power is equal to that of an asymmetric molecule containing four different radicals which are fairly short in length and have the values  $a$ ,  $b$ ,  $c$ , and  $d$  for their radii of repulsion.

The substitution  $AB = (a + b)$ , etc., in the determinants expresses the rotatory power completely in terms of the properties of the individual radicals. It is apparent that the expression must contain the differences  $(a - b)$ , etc., as factors, and probably the remaining factor will be a symmetrical function of almost constant value. It is convenient to resolve the determinants into alternants of the following type —

$$\begin{vmatrix} x \\ y \\ z \end{vmatrix} = \begin{vmatrix} L_1'' & L_2'' & L_3'' \\ L_1' & L_2' & L_3' \\ L_1'' & L_4'' & L_2'' \end{vmatrix} \quad \text{where } \begin{array}{l} L_1 = AB \quad CD \\ L_2 = BC \quad AD \\ L_3 = AC \quad BD. \end{array}$$

The advantage of this process is that every alternant can be resolved into the

\* 'Ann. Reps. Chem. Soc.', vol. 28, p. 385 (1931)



product of a symmetrical function and the simplest alternant, and this last factor is equal to the difference product

$$\begin{vmatrix} 0 \\ 1 \\ 2 \end{vmatrix} = (L_1 - L_2)(L_2 - L_3)(L_3 - L_1) \\ = -(a - b)(a - c)(a - d) \\ \times (b - c)(b - d) \\ \times (c - d) = -W \quad (28)$$

The tetrahedron which is being considered is of a special class, since

$$AB + CD = BC + AD = AC + BD = a + b + c + d = K,$$

and this relation enables the determinants of (27) to be resolved into alternants in a fairly straightforward manner. The general method is to separate the determinant so that symmetrical functions can be removed from a whole row leaving each element in terms of  $L_1$ ,  $L_2$ , and  $L_3$ . The resolution of the first and second determinants is carried out as follows —

$$\begin{aligned} \text{(I)} \quad & \begin{vmatrix} 1 & & \\ AB^5 & CD^4 + AB^4 & CD^5 \\ & AB^5 & CD^5 \end{vmatrix} = \begin{vmatrix} 1 & & \\ L_1^4(AB + CD) & & \\ & L_1^5 & \end{vmatrix} = K \begin{vmatrix} 0 \\ 4 \\ 5 \end{vmatrix} \\ \text{(II)} \quad & -2 \begin{vmatrix} 1 & & \\ L_1^3(AB^5 + CD^5) & & \\ & L_1^5 & \end{vmatrix} = -2 \begin{vmatrix} 1 & & \\ L_1^3(K^5 - 5K^3L_1 + 5KL_1) & & \\ & L_1^5 & \end{vmatrix} \\ & = -2K^5 \begin{vmatrix} 0 \\ 2 \\ 5 \end{vmatrix} + 10K^3 \begin{vmatrix} 0 \\ 3 \\ 5 \end{vmatrix} - 10K \begin{vmatrix} 0 \\ 4 \\ 5 \end{vmatrix} \quad (29) \end{aligned}$$

The resolution of the other determinants follows a similar course, though the last determinant is rather more complicated. It is necessary to use the following symmetrical functions —

$$\left. \begin{aligned} p &= \Sigma a, & K &= AB + CD \\ q &= \Sigma ab, & P &= \Sigma L_1 \\ r &= \Sigma abc, & Q &= \Sigma L_1 L_2 \\ s &= abcd, & R &= L_1 L_2 L_3 \end{aligned} \right\} \quad (30)$$

These are related and we have

$$\left. \begin{aligned} K &= p, & P &= 2q \\ Q &= q^2 + pr - 4s, & R &= pqr - r^2 - ps \end{aligned} \right\} \quad (31)$$

After complete resolution I is given by

$$\begin{aligned} I = 2KVR^{-1} & \left\{ -12 \begin{vmatrix} 0 \\ 4 \\ 5 \end{vmatrix} + (10K^2 + 5P) \begin{vmatrix} 0 \\ 3 \\ 5 \end{vmatrix} \right. \\ & + (2K^2P - 16Q + 2P^2 + 8s) \begin{vmatrix} 0 \\ 2 \\ 5 \end{vmatrix} \\ & + (10K^2Q - 6K^2P^2 + 5PQ - 5R - 24K^2s) \begin{vmatrix} 0 \\ 1 \\ 5 \end{vmatrix} \\ & \left. + 3R \begin{vmatrix} 0 \\ 2 \\ 4 \end{vmatrix} + (3PR - 12K^2R) \begin{vmatrix} 0 \\ 1 \\ 4 \end{vmatrix} \right\} \quad (32) \end{aligned}$$

All alternants can be resolved into a product of the lowest alternant and a symmetrical function of the same variables. Thus by the ordinary methods of manipulation of determinants we have the following identities —

$$\begin{aligned} \begin{vmatrix} 0 \\ 1 \\ 4 \end{vmatrix} &= (P^2 - Q)(-W), & \begin{vmatrix} 0 \\ 2 \\ 4 \end{vmatrix} &= (PQ - R)(-W), \\ \begin{vmatrix} 0 \\ 1 \\ 5 \end{vmatrix} &= (P^2 - 2PQ + R)(-W), & \begin{vmatrix} 0 \\ 2 \\ 5 \end{vmatrix} &= (P^2Q - Q^2 - PR)(-W), \\ \begin{vmatrix} 0 \\ 3 \\ 5 \end{vmatrix} &= (PQ^2 - QR - P^2R)(-W), & \begin{vmatrix} 0 \\ 4 \\ 5 \end{vmatrix} &= (Q^2 - 2PQR + R^2)(-W) \end{aligned} \quad (33)$$

Substituting these in (21)

$$\begin{aligned} I = 2KWVR^{-1} (6K^2P^2 - 24K^2P^2Q + 24K^2P^2s + 30K^2P^2R \\ + 12K^2PQ^2 - 48KPQs - 12K^2QR + 24K^2sR \\ - 7P^4Q + 9P^3R + 23P^2Q^2 - 8P^2Qs - 50PQR + 8PsR - 4Q^3 \\ + 8Q^2s + 20R^2) \end{aligned} \quad (34)$$

At the present stage it would be quite possible to evaluate the specific rotatory powers of actual compounds, but the evaluation of the symmetrical function  $S$ , contained in the brackets, for each compound would be very laborious. The value of  $S$  is not very sensitive to the differences of  $a$ ,  $b$ , etc., and an approximation can be made by replacing  $a$ ,  $b$ ,  $c$ , and  $d$  by the mean value  $(a + b + c + d)/4$ , i.e.,  $K/4$ . In the case of amyl alcohol this involves an error of 10%. We can make a more accurate approximation by a substitution of the following symmetrical functions for those in (34) retaining  $K$

$$\begin{aligned} \Sigma (3a - b - c - d)^2 / (a + b + c + d)^2 &= \Sigma (4a - K)^2 / K^2 = \Delta_2 \\ \Sigma (4a - K)^3 / K^3 &= \Delta_3 \\ \Sigma (4a - K)^4 / K^4 &= \Delta_4 \end{aligned} \quad (35)$$

The result is a series such as

$$S = AK^{12} (1 + A_1\Delta_1 + A_2\Delta_2 + A_3\Delta_3 + A_{22}\Delta_2^2 + A_{23}\Delta_2\Delta_3 \dots),$$

where the first term  $AK^{12}$  is the first approximation, obtained when  $K/4$  is substituted for  $a$ ,  $b$ ,  $c$ , and  $d$ , and the higher terms are purely difference terms giving a more accurate value of  $S$ . The value of  $\Delta_4$  and  $\Delta_2\Delta_3$  terms are very small and accordingly all higher terms will be neglected.

The substitution of  $\Delta_2$ , etc., in (34) is carried out by the identities

$$\left. \begin{aligned} 16q/K^2 &= 6 - \frac{1}{2}\Delta_2 \\ 64r/K^3 &= 4 - \Delta_2 + \frac{1}{2}\Delta_3 \\ 256s/K^4 &= 1 - \Delta_2 + \frac{1}{2}\Delta_2 - \frac{1}{4}\Delta_4 + \frac{1}{8}\Delta_2^2 \end{aligned} \right\} \quad (36)$$

and

$$\left. \begin{aligned} P/K^2 &= \frac{1}{2} (1 - \frac{1}{4}\Delta_2) \\ Q/K^4 &= \frac{1}{16} (1 - \frac{1}{2}\Delta_2 + \frac{1}{8}\Delta_4 - \frac{1}{16}\Delta_2^2) \\ R/K^6 &= \frac{1}{64} (1 - \frac{1}{2}\Delta_2 + \frac{1}{16}\Delta_4 + \frac{1}{8}\Delta_2^2 - \frac{1}{16}\Delta_2\Delta_3) \end{aligned} \right\} \quad (37)$$

These values have been substituted in each term of  $S$  and by addition of these expressions, while all terms higher than  $\Delta_2\Delta_3$  are neglected, we obtain

$$\begin{aligned} S = K^{12} 16^{-2} (207.5 - 92.25\Delta_2 + 17.5\Delta_3 - 51.9\Delta_4 + 42.5\Delta_2^2 \\ - 4.2\Delta_2\Delta_3) \end{aligned} \quad (38)$$

This method of expression, which is valuable when the values of  $a, b, c$ , and  $d$  are very close together, can be extended further to  $V$  and  $R^{-5}$ , both of which are symmetrical functions. These can be obtained in similar series and combined with  $S$  so that only one function need be evaluated. We have by the binomial theorem

$$R^{-5} = K^{-30} 2^{+30} (1 + 1.25 \Delta_2 - 0.3125 \Delta_4 + 1.015 \Delta_2^2) \quad (39)$$

The volume of a tetrahedron of edges  $(a+b)$ , etc., is derived from the expression for the volume when the three angles at an apex are  $\lambda, \mu$ , and  $\nu$  and the lengths of the edges meeting at this apex are  $x, y$ , and  $z$

$$\begin{aligned} V &= \frac{1}{6}xyz \sqrt{(1 - \cos^2 \lambda - \cos^2 \mu - \cos^2 \nu + 2 \cos \lambda \cos \mu \cos \nu)} \\ &= \frac{1}{6} \sqrt{(2 \sum a^2 b^2 c^2 d - \sum a^2 b^2 c^2)} = \frac{1}{6} \sqrt{(4qs - r^2)} \\ &= K^3 (\sqrt{2/96}) (1 - 0.375 \Delta_2 + 0.333 \Delta_3 - 0.375 \Delta_4 + 0.117 \Delta_2^2 \\ &\quad + 0.125 \Delta_2 \Delta_3) \end{aligned} \quad (40)$$

Substituting values of  $S, V$ , and  $R^{-5}$  in (34)

$$\begin{aligned} I &= \frac{W}{K^{14}} \frac{2.2075 \sqrt{2} 2^{30}}{16^3 96} \\ &\quad \times (1 + 0.430 \Delta_2 + 0.418 \Delta_3 - 0.938 \Delta_4 + 0.480 \Delta_2^2 + 0.447 \Delta_2 \Delta_3) \end{aligned} \quad (41)$$

We could easily obtain the final rotation by use of this value, but by means of a small approximation the value of the series can be tabulated. We will substitute by the identity  $\Delta_2 \Delta_3 = \frac{1}{2} \Delta_5$  and make the approximation  $\Delta_2^2 = 4 \Delta_4$ . This is true when  $\Delta a^2 = \Delta b^2 = \Delta c^2 = \Delta d^2$ , but is just an approximation in the ordinary case.

Defining  $\Delta a = (4a - K)/K$  we can split the series in (41) into separate series in powers of  $\Delta a$ , etc

$$f(\Delta a) = 0.430 \Delta a^2 + 0.418 \Delta a^3 + 0.932 \Delta a^4 + 0.537 \Delta a^5 \quad (42)$$

The value of  $f(\Delta a)$  has been tabulated and is obtained from the value of  $4a/K$  by the use of Table I. The values of  $f(\Delta b)$ , etc., are obtained similarly from the corresponding values of  $4b/K$ , etc. The value of the series is obtained by adding these quantities to unity

$$I = C_1 W K^{-14} (1 + f(\Delta a) + f(\Delta b) + f(\Delta c) + f(\Delta d)) = C_1 W K^{-14} (1 + F)$$

In an extreme case ( $e.g., 4a/K = 1.6$ ) the quantities  $\Delta a$ , etc., might be in such a relation that errors up to 10% would be introduced, but for actual

compounds the error involved in using the table instead of the exact equation (41) will be much less

We are now in a position to put down the formula for the specific rotatory power of the classical optically active compound by putting I in equation (26)

$$[\alpha] = \frac{C(\mu^2+2)(\mu^2+5)R_A R_B R_C R_D (1+F)(a-b)(a-c)(a-d)(b-c)(b-d)(c-d)}{\lambda^2 M (a+b+c+d)^{14}} \quad (43)$$

$\mu$  is the refractive index of the medium in which the rotation is measured

Table I

$4a/K$	$f(\Delta a)$	$4a/K$	$f(\Delta a)$	$4a/K$	$f(\Delta a)$
0 40	0 150	0 60	0 062	1 44	0 165
0 42	0 139	0 70	0 034	1 46	0 187
0 44	0 129	0 80	0 015	1 48	0 211
0 46	0 119	0 90	0 004	1 50	0 238
0 48	0 109	1 00	0 000	1 52	0 267
0 50	0 100	1 10	0 005	1 54	0 300
0 52	0 091	1 20	0 022	1 56	0 335
0 54	0 084	1 30	0 059	1 58	0 373
0 56	0 075	1 40	0 128	1 60	0 414
0 58	0 069	1 42	0 145		

$R_A$ , etc., are the total refractivities of each attached group, as given in the tables of Bruhl or Eisenlohr

$a, b, c$ , and  $d$  are the effective radii of the groups in Angstrom units

$f(\Delta a)$ , etc., have not much effect on the numerical result and, after finding  $4a/(a+b+c+d)$  are obtained from Table I

$\lambda$  is the wave-length of light used.

$M$  is the molecular weight of the compound

$C$  is a constant

$$C = \frac{72900 \cdot 207 \cdot 5 \cdot \sqrt{2} \cdot 2^{36} \cdot 2 \cdot 10^{64}}{32 \pi^2 16^3 96 N^3} = 16 \cdot 62,$$

taking the value  $N = 6 \cdot 06 \times 10^{23}$  and introducing  $10^{64}$  in order that the lengths of the tetrahedron might be expressed in Angstrom units

For light of D lines  $C/\lambda^3 = 4 \cdot 786 \times 10^9$ , and for this wave-length this is conveniently used as a constant

$[\alpha]$  is the specific rotation for a molecule of such configuration that, when the group A is placed nearest to the observer, the groups B, C, D appear in clockwise order. This gives the interesting result that the actual spatial distribution is known and it is no longer necessary to refer the spatial distribution of

most compounds to some reference compound. According to this analysis, we can say with absolute certainty that a dextro-compound has the configuration such that, when the largest group is nearest to the hypothetical observer, the other groups in order of diminishing size appear in a clockwise rotation. It may be noted that this result depends on taking the positive sign for the square root in the evaluation of  $V$  by equation (31), this is essential as  $V$  was defined as  $\frac{1}{8}[\text{AB AD CD}]$  which is positive for a tetrahedron of the configuration described above.  $[\alpha]$  was also taken as a rotation in the clockwise direction of the plane of polarization as seen by the observer at the receiving end of the polarimeter, this is the accepted convention for terming compounds dextro, apart from such compounds as the sugars where the prefixes are used to denote the relation of the configuration to that of some reference compound. The difference product has been put in the form  $(a-b)(a-c)(a-d)(b-c)(b-d)(c-d)$  so that this is found to be positive when  $a > b > c > d$ .

The final formula has now been derived and this gives the rotatory power of a compound containing four different radicals in terms of their properties, but only when the radicals are small. If we have data giving the absolute magnitudes of  $R_A$ ,  $\alpha$ , etc., we can calculate the rotatory power of some compounds of this type, such as  $\text{C}_2\text{H}_5\text{CHOHCH}_3$ . The evaluation of  $R_A$  and  $\alpha$  must be examined to see how these depend on the atoms in the radical.

The refractivity of a radical is obtained directly by the addition of the atomic refractivities of its constituents, but the effect of the central atom must also be considered. The valency electrons of this atom will act almost as part of the attached radicals and move with these to take up a dissymmetric distribution. If the central atom is small, as in carbon, since most of its refractivity is due to these electrons we can regard it as completely shared by the larger radicals, and we shall correct the refractivities of the radicals by adding one-quarter of that of the central atom.

The quantities  $a$ , etc., were termed the radii of repulsion of the radicals, but rather than representing any exact physical quantity they must be regarded more as parameters used to express the shape of the molecule. The radical is usually more complex than a single atom. In a single atom we might have regarded the radical as having a radius which could not be penetrated equal to that of the orbit of the outer electrons. In several atoms joined together our knowledge of the repulsion effects is very vague and the simplest assumption is that the total volume guarded from penetration is equal to the sum of the atomic volumes. The atomic radii are obtained by finding the distance

between atomic nuclei in diatomic molecules, and taking the total distance as the sum of the radii of the two atoms concerned. It will be noted that the radius here has very much the same meaning as that in which we have employed it in the above treatment. The measurements of the molecular dimensions are mainly from rotation spectra, and have been confirmed by X-ray and electron diffraction of vapours. The atoms are considered as spheres to obtain their volumes and then the radical assumed spherical to deduce the radius from the total volume. The radical radius is thus the cube root of the sum of the cubes of the atomic radii. As with the refractivity we shall correct for the effect of the central atom by adding one-quarter of its volume, this should give the approximate effect since the volume of the central atom will prevent the radicals approaching when they are small, but will have little effect when the radicals are large.

We can represent the method of calculating  $R$  and  $a$  by

$$R_A = \Sigma_A R + R(c)/4 \quad \text{and} \quad a = (\Sigma_A r^3 + r(c)^3/4),$$

where  $R(c)$  and  $r(c)$  are refractivity and radius of the central atom.

In the following calculation the refractivities for the D lines according to Eisenlohr are taken from Landolt-Bornstein 'Tabellen,' 1923 edition. The atomic radii H 0.37 Å, C 0.77 Å, O 0.70 Å, and N 0.72 Å are taken from 'Ann. Reps. Chem. Soc.,' vol. 28, p. 402 (1931). Table II, data from various sources, chiefly rotational spectra. In accordance with the suggestion in the Report, the radii for O and N have been corrected on the assumption that double and triple bonds cause 14% and 23% shortenings in length since the radii are taken from the molecules  $O_2$  and  $N_2$ . The properties of the radicals  $C_2H_5$ ,  $CH_3$ ,  $OH$ , and  $H$  have been found and substituted in formula (43) to obtain the rotatory power of sec butyl alcohol, and the other calculations performed in a similar manner. The values of the refractive indices have been taken from Landolt-Bornstein 'Tabellen.'

The experimental data are from sources commonly quoted for these compounds.

The concordance between the experimental and calculated values is better than the accuracy of the original data and the method of calculation warrants. This close agreement must be fortuitous, and the points of main importance are that the rotations forecasted are of the correct order and that the ratios of the values in the four compounds considered are very much the same in both the theoretical and experimental series. The four compounds chosen are the simplest active compounds, but they give a valuable proof that formula

(43) will apply for the calculation of the rotation of compounds in which we can regard the radicals as approximately spherical. This is the first time that the absolute magnitudes of the rotatory powers of actual compounds have been calculated from data entirely independent of optical activity measurements.

Table II—Specific Rotatory Powers  $[\alpha]_D$ 

	Calculated	Experimental	Observer
Amylamine	3.6	5.86	Marckwald, 'Ber. deuts. chem. Ges.', vol. 37, p. 1045 (1904)
Amyl alcohol	4.0	5.90	Marckwald and McKenzie, <i>ibid.</i> , vol. 34, p. 490 (1901)
Sec. butylamine	7.4	7.44	Thomé, <i>ibid.</i> , vol. 36, p. 586 (1903)
Sec. butyl alcohol	9.3	13.9	Pickard and Kenyon, 'J. Chem. Soc.', vol. 99, p. 45 (1911)

While we have employed formula (43) to forecast the absolute rotations from atomic data it is not in this direction that it will be most useful. The method of obtaining the radical radii from atomic radii, uncertain quantities in themselves, precludes the application to more complex compounds, but having established formula (43) it is possible for us to use it to obtain relative rotations. It seems to me that in the parachor we have a reliable method of obtaining an estimation of radical radius, the more so because this will already include any special repulsion effects. To use the cube root of the parachor for quantitative predictions it would be necessary to obtain an empirical constant for the formula, but the idea may be useful for the qualitative study of the variations of rotation by substitution.

In a large radical it appears that only the portion adjacent to the asymmetric centre will enter into the rotatory power interactions, and the only effect of those portions of the radical far removed from the asymmetric centre will be to repel the adjacent radicals slightly, thus substitution at the far end of a chain will only alter the rotatory power to the extent that it alters the configuration around the centre. Substitution of a group with a large parachor should increase the effective radius of the group which it enters, and, if the relative sizes of the groups are known previously, it should be possible to forecast the direction of the change in rotation. The effect will be much greater if the substitution is near the asymmetric centre.

An examination of the parachors, especially of those portions near the asymmetric centre, should provide a means of estimating the absolute con-



figuration of any enantiomorph from the sense of its rotatory power. This can only be done with certainty when interactions of the solvent with the molecule or association are known to be absent, but, when a reliable estimation can be made, this should be of vital importance in the study of Walden Inversion reactions.

In formula (43) we have the connecting link between rotation and chemical structure. Although the formula is only applicable in a quantitative manner to the extent that the molecule approximates to four spherical groups, it demonstrates clearly what variations in rotation are to be expected for known changes in the atomic structure.

#### *Rotatory Power, Rotatory Dispersion, and Molecular Structure*

It is definitely established that, if a compound containing four groups attached to a central atom exhibits optical activity, all these groups must be different chemically. Since the very existence of a rotation depends on the differences of these groups, it appears that the magnitude of this rotation must depend on the difference of some specific property  $A$  of the groups, and that there must be a factor of the type  $(A - B)(A - C)(A - D)(B - C)(B - D)(C - D)$  in any theoretical expression for the optical rotation. This logical deduction was made independently by Crum Brown\* and Guye,† but these investigators had no definite idea of the mechanisms involved in the actual twisting of the plane of polarization. Crum Brown also showed that, if one group could be changed in small increments so that the magnitude of this property in it approached the magnitude of this property in another group, the rotation would decrease to zero and then change sign as the first group surpassed the fixed group.

It has been established in this paper that the rotatory power is given by a formula of the predicted kind, in which the effective radius is the property occurring in the continued difference product. If one group can be changed while the compound retains the same configuration, the sign of the rotation should change when the effective radius is increased above that of the next largest group. This has been shown experimentally if we can assume the effective radius of an aliphatic chain to increase with its length, Kuhn‡ has observed that in the series of compounds  $R_1 R_2 \text{CHOH}$ , if these are prepared in a manner such that the configuration is definite, then the sign of the rotation

\* 'Proc. Roy. Soc., Edin.,' vol. 17, p. 181 (1890).

† 'C. R. Acad. Sci., Paris,' vol. 110, p. 714 (1890).

‡ 'Trans. Faraday Soc.,' vol. 22, p. 457 (1926).

is constant if  $R_1 > R_2$ . Again in active amyl derivatives we find that  $-\text{OH}$  and  $-\text{NH}_2$  are the only radicals which have less atomic volumes than the  $-\text{CH}_3$  group, all other radicals cause the  $-\text{CH}_2\text{X}$  group to be larger than the adjacent  $-\text{CH}_2\text{CH}_3$  group. Thus amyl alcohol and amylamine should have the opposite sense of rotation to all other compounds prepared by substituting for  $-\text{OH}$  in amyl alcohol, and this is entirely fulfilled experimentally.

It is interesting to note that we can ascertain the absolute configuration of the ordinary amyl alcohol, which is levorotatory. This is given in fig 1, when the  $\text{CH}_2\text{OH}$  group is placed nearest to the observer the other groups taken in clockwise order are  $\text{C}_2\text{H}_5$ ,  $\text{CH}_3$ , and  $\text{H}$ .

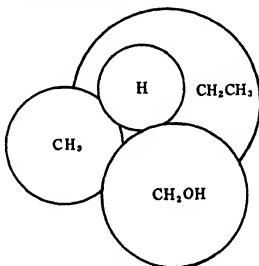


FIG 1—Absolute configuration of levorotatory amyl alcohol. For simplicity the radicals have been represented as spheres.

An examination of formula (43) shows that it divides quite naturally into three factors which seem almost independent, and the variations of the rotatory power under different conditions are easily classified by considering the changes in these three factors  $(\mu^2 + 2)(\mu^2 + 5)$  dependent solely on the medium;  $R_A R_B R_C R_D / \lambda^2$  dependent on the refractivities and the wavelength, and

$$(1 + F)(a - b)(a - c)(a - d)(b - c)(b - d)(c - d)/(a + b + c + d)^4$$

totally dependent on the spatial arrangement

The first factor  $(\mu^2 + 2)(\mu^2 + 5)$  is more a property of the medium than of the active compound. In accurate work this factor should be corrected

for in order to obtain comparative results when working in different solvents and with different wave-lengths, but as a general rule the variations in the rotatory power brought about by this factor are very small

The second factor determines the variation in rotation with changing wave-lengths of light. In this examination the first factor will be neglected, or can be regarded as corrected for by suitable measurements on the refractive index of the medium at different wave-lengths. For many compounds there are regions in which  $R_A$ , etc., are almost indifferent to changes in wave-length, in these regions the rotatory dispersion is given by Biot's inverse square law. For other regions, since the refractivities  $R_A$ , etc., can be determined from ordinary refractive data, it is possible to predict exactly the rotatory dispersion. However, it is interesting to find the general form of the rotatory dispersion curve, which can be obtained since the general equation for refractive dispersion is known. For the sake of simplicity, the term corresponding to a dissipation of energy by absorption will be omitted and hence the resulting rotatory dispersion equation will only apply for wave-lengths outside absorption bands. The extra terms can be introduced if desired.

According to the Drude treatment of refractivity as modified by Lorentz the refractivity of any group is obtained by summing for the motion of all its electrons,

$$3R = \frac{3(n^2 - 1)}{n^2 + 2} = \frac{3(K - 1)}{K + 2} + \frac{1}{4\pi c} \sum \frac{v_e e_e \lambda_e^4}{m_e (\lambda^2 - \lambda_e^2)}$$

The first term represents the refractivity for infinitely long waves and the second the sum of the effects of absorption bands situated at  $\lambda_e$ , etc. For regions not very near an absorption band only the first term is large, and, as it is not very probable that two absorption bands in different groups will be at the same wave-length, we may evaluate  $R_A R_B R_C R_D / \lambda^2$  neglecting portions which contain products of absorption terms

$$\frac{R_A R_B R_C R_D}{\lambda^2} = \frac{1}{\lambda^2} \left[ R_A' R_B' R_C' R_D' + R_A' R_B' R_C' \frac{K_D}{\lambda^2 - \lambda_D^2} + R_B' R_C' R_D' \frac{K_A}{\lambda^2 - \lambda_A^2} \right]$$

and

$$[\alpha] = \frac{1}{\lambda^2} \left[ K_0 + \frac{K_1}{\lambda^2 - \lambda_1^2} + \frac{K_2}{\lambda^2 - \lambda_2^2} \right] \quad (44)$$

Only one absorption band has been written for each group, but it is obvious that, if there are more than one, all these must be included. The last equation is the most general form of the rotatory dispersion equation, where  $\lambda_1$ ,  $\lambda_2$ , etc., are the wave-lengths at which there are absorption bands in the separate groups adjacent to the asymmetric centre

The form of equation which is generally used for expressing experimental results is that deduced by Drude on his hypothesis of spiral movements of electrons, and this equation has been well established by Lowry and his co-workers. Though the Drude form does not appear the same as the above equation, these can be directly transformed into each other. Taking the two terms which Lowry finds sufficient to represent the behaviour of most compounds we have

$$[\alpha] = \frac{K_1}{\lambda^2 - \lambda_1^2} + \frac{K_2}{\lambda^2 - \lambda_2^2}$$

$$= \frac{1}{\lambda^2} \left[ (K_1 + K_2) + \frac{\lambda_1^2 K_1}{\lambda^2 - \lambda_1^2} + \frac{\lambda_2^2 K_2}{\lambda^2 - \lambda_2^2} \right]$$

Since  $\lambda_1^2 K_1$  and  $\lambda_2^2 K_2$  are constants, this last equation has the same form as (44). The formula for the rotatory dispersion obtained on the assumption that the rotatory power of the molecule is due to the electric interactions of the induced electric doublets is just the same as that used in experiment and derived from other hypotheses, but it has the advantage that the significance of the constants are known exactly and their values can be derived independently from other sources.

The Drude formula for the rotatory dispersion resolves the rotatory power into the sum of several terms, each of which appears to be the contribution of a separate group of electrons giving one or more definite absorption bands. Occasionally a single term gives a very large contribution and it would appear that the electrons have a very dissymmetric structure. For camphor the rotatory power in the ultra-violet region is dependent largely on a term  $K/(\lambda^2 - \lambda_a^2)$  where  $\lambda_a$  is the absorption wave-length of the ketone CO group, but it seems impossible for a symmetrical group such as the ketone group to have a dissymmetric structure. In order to meet this difficulty Lowry and Walker\* formulated the interesting hypothesis that some influence could emanate from the asymmetric centre and induce asymmetry in adjacent groups such as the CO group. However, in this analysis we do not regard the rotatory power as due to asymmetry concentrated in various centres, and this phenomenon should be completely explained according to the factor  $R_A R_B R_C R_D / \lambda^2$ .

Considering the region just outside the absorption band we see that  $R_{CO}$ , the refractivity of the ketone group, will vary very rapidly with  $\lambda$ . This variation will be so great that, in comparison, we can regard  $(R_A R_B R_C)$  as

\* 'Nature,' vol. 113, p. 565 (1924).

practically constant The ketone group is considered as the fourth refractive centre  $D$  and the remaining factor  $R_{\lambda 0}/\lambda^3$  is expressed by  $K/(\lambda^2 - \lambda_{\lambda 0}^2)$  according to the ordinary theory of refractive dispersion  $\lambda_{\lambda 0}$  is the absorption wave-length of the ketone group and so the variation of the whole rotatory power is given by a single Drude term dependent on this wave-length This is just the experimental result and there is no necessity for any further assumptions

In the last factor

$$(1 + F)(a - b)(a - c)(a - d)(b - c)(b - d)(c - d)/(a + b + c + d)^4$$

we have an expression which is characteristic of a formula for optical activity The exact predictions and applications of this have already been discussed, but it is noteworthy that since this factor has a very differential nature most of the variation of activity from compound to compound is given by this factor The variations of  $R_A$ , etc., in different compounds are small and we may regard the spatial configuration as altogether the most important property in the determination of the rotatory power

The quantities  $a$ ,  $b$ ,  $c$ , and  $d$  were only given a meaning according to the Thorpe-Ingold theory of valency deflexion Since the numerical calculations for the simple compounds *sec*-butyl alcohol, etc., agree with the experimental data, this constitutes another point of evidence in favour of the Thorpe-Ingold theory

Hitherto we have ignored the occurrence of permanent dipoles in certain molecules A permanent dipole will not interfere with the electric interactions causing the rotatory power, but it may have a definite effect on the configuration A polar group may tend either to attract or repel the adjacent groups according to their nature We can see the influence of a polar group most clearly if we consider it at such a distance that its volume does not affect the configuration A dipole at the end of a long chain may still have an appreciable deforming influence on the portions around the asymmetric centre, the effect may be transmitted either as an electric field acting through space or as a polarization along the atoms linking with the centre We should expect the rotatory power to vary with the size of the dipole Rule\* and his collaborators have studied several series of compounds with the dipoles at a distance from the centre, and it is probable that the alteration of configuration around the asymmetric centre explains the systematic variation in rotatory power which has been found.

\* Rule and Smith, 'J. Chem. Soc.', vol. 127, p. 2188 (1925).

In the experimental study of the dependence of the optical activity on chemical constitution, one of the great difficulties has been the sensitivity of the rotation of many compounds to changes in solvent and temperature. The present analysis seems to provide a logical basis by which one can understand and correlate these variations with other properties of the compounds and of the solvents. The optically active compounds will fall roughly into three classes according to their atomic structure and these will give more or less distinct behaviours on variation of the physical conditions.

When we consider a molecule with little or no polar groups and no groups which have a tendency to co-ordinate with any other group, even if this is dissolved in a strongly polar solvent, it cannot be appreciably deformed by the internal electric fields. The refractivities will be affected much less, and the only variation will be due to the factor  $(\mu^2 + 2)(\mu^2 + 5)$ . Thus, for changes from one solvent to another and for changes of concentration while in one solvent, the specific rotation should vary as  $(\mu^2 + 2)(\mu^2 + 5)$  where  $\mu$  is the refractive index of the solution under exactly the same conditions as in the measurement of the rotation.

In a molecule containing a dipole we should expect this, by virtue of its local electric field, to exert a deforming influence in the molecule. The electric field will be partially transmitted through the space external to the molecule and its magnitude will be very dependent on the presence or absence of other polar molecules which will determine a mean dielectric constant of the external space. Thus in changing from solvent to solvent probably the variation of the specific rotation with  $(\mu^2 + 2)(\mu^2 + 5)$  will be entirely masked by variations dependent on the dielectric constant. Thus for inert polar molecules the variation of the specific rotation should give exactly the same serial order as arranging the solvents in order of their dielectric constants. In this and the preceding cases the change of medium will not alter the chemical nature of any of the groups, and we should not expect any change in the form of the rotatory dispersion curves.

In the molecule which contains a group which can co-ordinate with other molecules, whether of the same or a different kind, the system is very complicated. In solution, or even in the pure state when the compound associates, the system will be an equilibrium consisting, as regards optical activity, of entirely different molecules. The group that co-ordinates will have its effective radius very much increased by junction with another whole molecule and the magnitude and sign of the rotation may be altogether altered. The direction of this change may be predicted if we know from chemical evidence

which group it is that co-ordinates. However, in the loose linking with another molecule the chemical structure of the group may be so altered that the refractivity and refractive dispersion are changed. Whereas all other types of compound retain the same rotatory dispersion curve even if the magnitude of the rotation is changed by concentration or solvent, this type of molecule which co-ordinates with another molecule may have both the rotation and rotatory dispersion entirely altered.

There is no direct effect of the temperature apparent in the general case and the variations due to temperature are due to variations in the properties of the medium, *e.g.*, refractive index, dielectric constant, and equilibrium in association. From the theoretical viewpoint the temperature and solvent action can be classified according to nature of the compound. (1) The compound is inert and the rotation depends on the refractive index of the medium while the rotatory dispersion is constant. (2) The compound is polar, but inert, and the rotation depends chiefly on the dielectric constant while the rotatory dispersion is constant. (3) The compound is active chemically, tending to co-ordinate, and both rotation and rotatory dispersion vary with temperature and concentration.

With regard to the relation of chemical constitution to rotatory power, rotatory dispersion, and solvent and temperature effects, the theoretical predictions seem to provide a clear understanding of the actions which lie behind these phenomena.

In conclusion, it may be emphasized that, since the direct calculation of the polarizations in the various refractive centres of the active molecule predicts rotations of the observed magnitudes, there is no necessity for any special molecular structure to be assumed as an explanation of optical activity. Indeed, since the actions which are the origin of the rotatory power are so well established by the electronic theory of dispersion, it appeared that this analysis could not be called an hypothesis, but could only be correctly named a calculation of optical activity.

I wish to thank the Rector of the Imperial College, H. T. Tizard, C.B., F.R.S., and Professor S. Chapman, F.R.S., for helpful suggestions in the presentation of these papers, and also Professor H. B. Baker, C.B.E., LL.D., F.R.S., for his encouragement and constant interest during the course of this work.

*Summary*

(1) The optical rotatory power of an asymmetric atom attached to four small radicals has been calculated and is expressed in a practical formula. The rotation is dependent on the refractivities and the sizes of the radicals.

(2) The formula has been confirmed by the calculation of the rotatory powers of amyl alcohol, amylamine, sec-butyl alcohol, and sec-butylamine.

(3) The conclusions drawn from this calculation with regard to optical activity and chemical substitution and optical activity and physical conditions agree with the characteristic behaviour of optically active compounds.

(4) An exact formula connecting the rotatory dispersion of the compound with the refractive dispersion of the radicals has been obtained.

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*Transmutation Effects Observed with Heavy Hydrogen*

By M. L. E. OLIPHANT, Ph.D. (Messel Research Fellow of the Royal Society),  
P. HARTECK, Ph.D., and LORD RUTHERFORD, O.M., F.R.S.

(Received April 14, 1934)

[PLATE 16]

In our paper "Transmutation of Elements by Protons,"\* we showed that the transformation of some of the light elements by protons could be conveniently studied by the use of comparatively low voltages—of the order of 100,000 volts—by generating an intense narrow beam of protons which fell on the target of small area of about 1 sq. cm. In the light of experience of the past year, the installation has been modified in several particulars and entirely reconstructed. By the addition of another 100,000-volt transformer in tandem and the use of appropriate condensers the D.C. voltage available has been raised from 200,000 to 400,000 volts. The main change, however, consists in the use of a horizontal instead of a vertical discharge tube. In place of glass, a corrugated porcelain wall bushing capable of withstanding high voltages has been used to insulate the positive electrode, while the earthed metal casing forming the negative electrode projects through a brick wall. The arrangement of the internal electrodes is, in general, similar to that used in the

\* 'Proc. Roy. Soc.,' A, vol. 141, p. 259 (1933).



earlier apparatus. The oil cooling circulation has been improved as the electrodes cannot now be cooled by radiation alone. As before a magnetic field is applied to sort out the various types of ions generated in the discharge tube. The use of the horizontal tube has many advantages, not only for assembling the controls at convenient points but also in the ease of handling the counting apparatus and absorbing screens.

The new installation has worked smoothly and satisfactorily and we have been able to increase the number of disintegration particles available for study by a factor of 10 to 50. The thick brick wall acts as a complete screen for the X-radiation generated in the system.

In our last paper\* we gave an account of the transformations produced in lithium by the ions of heavy hydrogen. The heavy water used for this purpose was generously presented to us by Professor G. N. Lewis. For our present experiments we have depended on a supply of concentrated heavy water prepared in the Cavendish Laboratory by Dr P. Harteck†. For preliminary requirements a weak concentration of diplogen‡ of about 12% was generally used. Stronger concentrations up to 30% mixture with helium§ were necessary in order to study the emission of neutrons and protons. The action of diplogens on diplogens was studied by observation of the effects produced when diplogens were used to bombard targets covered with a thin layer of a preparation containing heavy hydrogen. These were ammonium chloride, ammonium sulphate, and orthophosphoric acid in which the normal hydrogen had been largely replaced by diplogen. The method of preparation was very simple. A small quantity of the normal ammonium salt or the phosphoric pentoxide was added to an excess of heavy water. An equilibrium was at once established between the concentration of hydrogen and of diplogen in the compound and in the water,|| and if a drop of the solution was placed upon a warm iron target and allowed to evaporate a stable but non uniform layer of a salt containing diplogen was left behind. The  $\text{ND}_4\text{Cl}$  was also deposited upon the target in the form of a very thin and a uniform layer by means of sublimation, but this very property renders the target unstable and liable to disappear rapidly

\* 'Proc Roy Soc.' A, vol 141, p. 722 (1933)

† 'Proc Phys Soc Lond.', vol. 46, p. 277 (1934)

‡ In a discussion on Heavy Hydrogen before the Royal Society on December 14, 1933 (see 'Nature,' p. 955 (1933)), the names diplogen (D) for the new isotope of hydrogen and diplogon for its nucleus seemed to find favour. Also 'Proc Roy Soc.' A, vol 144, p. 1 (1934)

§ Cf. Oliphant, Kinsey, and Rutherford, 'Proc Roy Soc.' A, vol 141, p. 722 (1933)

|| Lewis, 'Proc Amer Chem Soc,' vol 55, p. 3502 (1933)

under bombardment The phosphoric acid remained as a liquid film over the surface of the target Exceedingly small quantities of the substances are required

*The Action of Diprotons on Diploons*

*The Emission of Charged Particles*—The most interesting and important reaction which we have observed is that of heavy hydrogen on heavy hydrogen itself Experiment has shown\* that diplogen is not appreciably affected by bombardment with  $\alpha$ -particles from polonium, and we have been unable to detect any specific action of protons on diplogen for energies up to 300,000 e-volts We were therefore surprised to find that on bombarding heavy hydrogen with diploons an enormous effect was produced Fig 4, Plate 16, shows a reproduction of portion of an oscillograph record obtained in our first experiment We assumed at first that this was an effect due to radiation passing through the counting chamber as previous experiments had shown that X-rays could produce just the result observed, but subsequent observation at much lower bombarding potentials showed that we were dealing in reality with a very large emission of protons Examples of an oscillograph record obtained under these conditions are given in figs 5 and 6, Plate 16 The original observations were made on  $\text{ND}_4\text{Cl}$ , but in order to establish that the effects observed came from the action of D on D and not from the nitrogen or chlorine, we bombarded targets of  $(\text{ND}_4)_2\text{SO}_4$  and of  $\text{D}_3\text{PO}_4$  The absorption curves obtained for the three substances are given in fig 1 The shape of these curves is due to the fact that protons gave too small a deflection in the oscillograph to be easily counted except over the last five centimetres of their path

It is evident from fig 1 that there are present in each case two very prominent groups of particles of ranges 14.3 and 1.6 cm. respectively Careful counting of the records established that the numbers of these particles were identical within the errors of measurement The maximum size of the deflections produced on the oscillograph record by the particles in each group indicated that they both consisted of singly charged particles On these data it is natural to assume that the particles are emitted in pairs opposite one another, and that the difference in range arises from a difference in mass, and hence of the velocity and energy The simplest reaction which we can assume is



\* Rutherford and Kempton, 'Proc. Roy Soc.' A, vol 143, p. 724 (1934)

† It has been decided to put the symbol for the nuclear charge to the left and below the symbol for the element

While from the point of view of both experiment and theory, we have no information of the details of the nuclear changes involved and can only study the final result, yet it is often useful and may even be more correct to assume that the first step in the process is the capture of the bombarding particle to form a new and heavier nucleus. If this proves unstable, it then breaks up, possibly in a variety of ways. The recent discovery by M and Mme Curie-Joliot,\* and by Cockcroft and Walton,† that a type of radioactive nucleus is

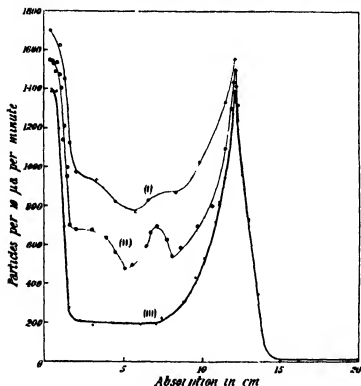


Fig 1—(i)  $\text{ND}_4\text{Cl}$ , (ii)  $(\text{ND}_4)_2\text{SO}_4$ , (iii)  $\text{D}_3\text{PO}_4$

formed by the capture of an  $\alpha$ -particle, dipion, or proton lends support to this point of view. The time of transformation may vary over very wide limits and may sometimes be so short that the process cannot be experimentally followed. In the case which we are considering we are inclined to interpret the observations in the following way. The initial process is the union of two dipions to form a new nucleus of charge 2 and mass 4, i.e., a helium nucleus

\* 'C. R. Acad Sci Paris,' vol 196, p. 254 (1934).

† 'Nature,' vol 133, p 328 (1934).

If we neglect the energy of the bombarding particle and assume the mass of D to be that given by Bainbridge,\* the mass of this helium atom must be 4.0272, and it therefore possesses an excess energy over the normal helium atom, of mass 4.0022, of about 23 million volts. This atom is unstable and may lose its energy in a variety of ways, some of which are considered later. We are considering here the breaking up of the helium nucleus into a proton and a hydrogen isotope of mass 3. The transformation follows so rapidly after the capture that no evidence of the existence of the excited helium nucleus has so far been obtained.

The mass and energy relations on the right-hand side of equation (1) can be obtained in the following way. The  ${}_1\text{H}^1$  particle possesses an energy of  $3.0 \times 10^6$  e-volts, corresponding to protons of the observed range of 14.3 cm. Then, from momentum considerations, the energy of the  ${}_1\text{H}^2$  particle which is emitted in the opposite direction will be  $1.0 \times 10^6$  e-volts, i.e., the total energy of the two particles will be  $4.0 \times 10^6$  e-volts, corresponding to a mass-change of 0.0043 units. Hence the mass of the  ${}_1\text{H}^2$  atom will be

$$4.0272 - (1.0078 + 0.0043),$$

i.e., 3.0151. The ionization produced by the  ${}_1\text{H}^2$  particle will be at every point of its path identical with that produced by a proton possessing the same velocity. However, owing to its greater momentum the  ${}_1\text{H}^2$  particle will travel three times as far as the proton for a given reduction of velocity. Consequently it will have three times the range of a proton of the same initial speed. The initial velocity of the  ${}_1\text{H}^2$ , corresponding to an energy of  $10^6$  e-volts, is  $8 \times 10^8$  cm/sec. The range of a proton with this velocity is 5.8 mm according to data given to us by Feather. Hence the range of the  ${}_1\text{H}^2$  particle should be  $3 \times 5.8 = 1.74$  cm. Considering the nature of the data available and the difficulties of determining the range of the short 1.6 cm. group accurately, we feel that this agreement is very satisfactory.

Additional evidence of the truth of our assumption is afforded by observation of the way in which the ionization, as measured from the magnitude of the oscillograph deflections, varies near the end of the range of the particles. In fig. 2 we have plotted the number of deflections above a given size against the absorption in the path of the particles, using a chamber 3 mm. deep. It is seen that the 14 cm. group shows a very sharp peak which occurs about 2 cm. short of the end of the range. The 1.6 cm. group, on the other hand, rises rapidly to what appears to be a much broader maximum. This is just what

\* 'Phys. Rev.', vol 44, p. 56 (1933).

would be expected from a particle of greater mass, the velocity of which varies less rapidly with the length of path in the material than the proton

Still further confirmation of the truth of this mode of transformation would be provided if it could be shown that the  ${}_1\text{H}^1$  and  ${}_1\text{H}^3$  particles recoil in opposite directions. This point has been carefully examined by Dee using the Wilson expansion chamber method, and in a recent letter to 'Nature'\* he concludes that there is no doubt of its correctness

It seems clear that the production of this isotope of hydrogen of mass 3 in these reactions is established beyond doubt. The mass of the  ${}_1\text{H}^3$  atom is consistent with its possessing a stability of the same order as  ${}_1\text{H}^3$ . The possible existence of this isotope has been discussed by several writers and although a

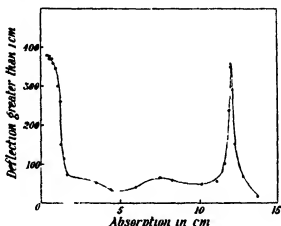


FIG 2— $(\text{ND}_4)_2\text{SO}_4$

careful search has been made no evidence of its presence has been found. It seems probable, however, that it could be formed by the process we have considered in sufficient quantity to be detected ultimately by spectroscopic or positive-ray methods.

**Voltage Variation and Absolute Yield**—The variation with energy of the bombarding deuterons, of the emission of 14.3 cm protons, has been measured over a limited range of voltage. The yield is so great that the number of particles entering the chamber soon became inconveniently large, and also we found that at high bombarding energies the heavy hydrogen compound is rapidly removed from the target. Fig 3 gives the curve obtained with  $(\text{ND}_4)_2\text{SO}_4$ . It is evident that particles are detected at energies as low as 20,000 volts

\* 'Nature,' vol. 133, p. 564, April 14th (1934)

and that the number increases rapidly with increase of accelerating potential\*. The nearly linear rise beyond 100,000 volts is probably due to the fact that, for potentials greater than this, the chance of disintegration in a collision is nearly constant, and the number of transformations therefore increases proportionally with the penetration of the bombarding particle into the target material.

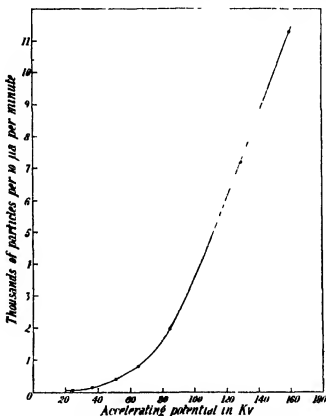


FIG. 3— $(\text{ND}_2)_2\text{SO}_4$ , 12 cm. absorber

From the solid angle subtended by the window, and the known composition of the target, we estimate that at 100,000 volts the absolute yield of disintegrations for collisions between D and D is of the order of magnitude of 1 in  $10^6$ . This is a far greater yield than that obtained for any other disintegration process, even at much higher potentials. Simple calculation shows that even a monomolecular layer of diplogen under the conditions of our

\* It should be emphasized that the energy available in a collision between two particles of equal mass is only half the energy of the bombarding particle, since half the kinetic energy is retained by the composite nucleus formed by capture.

experiment would give a large effect. Consequently every substance we have bombarded with diplogens begins, after a short time, to show effects which are clearly due to traces of diplogen absorbed by or driven into the target. We have also observed that when a compound containing hydrogen, such as ammonium sulphate or lithium hydroxide, is bombarded with a strong beam of diplogens, the hydrogen in the compound is gradually replaced in part by diplogen. After some time such compounds begin to behave as diplogen targets of appreciable concentration, and great care must be exercised to separate these effects from any others which are under investigation. This deposition of diplogen from the beam itself, while easily distinguished by the appearance of protons of characteristic range, is much more difficult to allow for when observations are made on the neutron emission of elements. For example, it is difficult to disentangle the emission of neutrons characteristic of lithium and beryllium from the spurious effect due to contamination by diplogen.

*The Emission of Neutrons*—We have pointed out that the appearance of the oscillograph records obtained in our initial experiments on the effects produced on bombarding diplogen with diplogen were suggestive of the presence of a strong radiation. The unstable form of He of mass 4.00272, formed by the union of two diplogens, might be able to revert to the normal form of He of mass 4.0022 by losing the additional mass as energy of a gamma ray or rays of  $23 \times 10^6$  e-volts energy. Accordingly we searched for such a radiation with a Geiger-Müller counter. It was at once evident that there was present a very intense radiation capable of producing an undiminished effect on the counter through 20 cm. of lead. As a check on this a search was made for recoil nuclei with the linear counter, and it was found that neutrons are emitted in numbers comparable with the number of 14 cm. protons. It is known that the Geiger counter is affected by neutrons both by the recoil nuclei produced in the counter itself, and also through the action of secondary radiations\* produced when neutrons pass through matter. Under these conditions it is impossible for us to decide, on the basis of our experiments, whether a gamma ray of high energy is present. In order to establish the existence of such a radiation it will be necessary to search for high speed photo-electrons, either with the expansion chamber or with a system of coincidence counters.

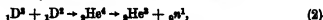
We have endeavoured to determine the properties of the neutrons by various methods. The absorption in lead has been measured with the linear counter

\* Cf. Lee, 'Nature,' vol 133, p 24 (1934). Experiments on these points have been made with the assistance of Mr Westcott, who provided the counters, and details will be published elsewhere.

by observation of the number of recoil kicks produced with or without an absorbing screen of 5.5 cm. of lead between the chamber and the target. The average of three measurements showed that the number of kicks was reduced by 44%, corresponding to a radius of cross-section for a collision between a neutron and a lead nucleus of  $8.4 \times 10^{-13}$  cm. This is to be compared with the value found by Chadwick for the neutrons from beryllium bombarded by polonium  $\alpha$ -particles, i.e.,  $7 \times 10^{-13}$  cm.\* Chadwick states that the cross-section does not appear to change rapidly with the energy of the neutron except when the energy is very low, so that the result tells us nothing except that the neutrons appear to behave in very much the same way as the neutrons from beryllium, which we know to possess a large range of energies.

It is possible to obtain an approximate value for the maximum energy of the neutrons by observation of the maximum size of the kicks produced by recoil nuclei in the linear counting chamber. From the known size of the kicks produced by  $\alpha$ -particles, and our knowledge of the energy loss of such particles in traversing the chamber, we can estimate the ionisation produced by, and hence the energy of the recoil nuclei. Thus we find that the maximum size of kick produced in the chamber by the recoil nuclei in air is about 6 mm., if we neglect the very large deflections of 2 cm. and over, which almost certainly result from disintegrations. Assuming that the energy loss of the recoil particle per ion produced along its path is the same as for an  $\alpha$ -particle, we find an energy for this nucleus of about  $0.5 \times 10^6$  e-volts. Application of momentum considerations to a head-on collision between a nitrogen nucleus and a neutron leads to the conclusion that the energy of the colliding neutron must be about  $2 \times 10^6$  e-volts. We have also made experiments on the energy of recoil nuclei produced in helium resulting in a slightly higher value of  $2.2 \times 10^6$  e-volts. This gas is especially suited for neutron recoil observations as it does not suffer disintegration by neutrons.

In order to account for the production of neutrons of the observed energy and number we have been led to assume the transformation



in which the unstable  ${}_2\text{He}^4$  nucleus first formed breaks up into a helium isotope of mass 3 and a neutron. The assumption of the formation of  ${}_2\text{He}^3$  as a product of such a transformation is not without a precedent, as we have already con-

\* 'Proc. Roy. Soc., A, vol. 136, p. 693 (1932).





FIG. 4. First observation of 14 cm protons from  $D + D$



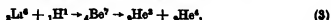
FIG. 5.—14 cm protons from  $D + D$  (12 cm absorber)



FIG. 6.  ${}^3\text{H}$  particles from  $D + D$  (1.1 cm absorber)



cluded that it is produced, together with a normal  $\alpha$ -particle, as a result of the bombardment of  ${}^6_3\text{Li}$  with protons.\* We assumed



and found that the observed ranges of 11.5 and 6.8 mm were in good accord with the application of momentum considerations to this reaction. Dees† has obtained very definite evidence that the two short-range particles are emitted in opposite directions, and we have confirmed that it is the  ${}^6_3\text{Li}$  isotope which is concerned in this transformation‡. It seems quite clear, therefore, that  ${}^3_2\text{He}$  can exist under these conditions, and from the observed ranges of the particles we are able to calculate its mass in the following way. The longer range of 11.5 mm is reasonably well known and it will correspond with the  ${}^4_2\text{He}$  particle. For the same velocity an  $\alpha$ -particle would go 4/3 times as far because of its greater energy. Hence the velocity of the  ${}^3_2\text{He}$  is the same as the velocity of an  $\alpha$ -particle the range of which is 1.53 cm., corresponding with an energy of  $2.5 \times 10^6$  e-volts. The energy of the  ${}^3_2\text{He}$  will be three-quarters of the energy of this  $\alpha$ -particle, i.e.,  $1.88 \times 10^6$  e-volts. The total energy produced in the transformation is thus  $3.30 \times 10^6$  e-volts, or the corresponding change in mass is 0.0035 units. Hence the mass of the  ${}^3_2\text{He}$  is

$$(6.0145 + 1.0078) - (4.0022 + 0.0035) = 3.0166,$$

using Bainbridge's (*loc. cit.*) value for the mass of  $\text{Li}^6$ . On the other hand, we can use the mass 6.0157 calculated from the disintegration of  ${}^6_3\text{Li}$  into two  $\alpha$ -particles of 13.2 cm. range under bombardment by deuterons,§ the data for this transformation being extremely good. From this transformation we obtain for  ${}^3_2\text{He}$  a mass of 3.0178 units.

Substitution in reaction (3) of the masses calculated in the above manner leads to the appearance of excess masses of

$$(2.0136 + 2.0136) - (3.0166 + 1.0067) = 0.0039,$$

and

$$(2.0136 + 2.0136) - (3.0178 + 1.0067) = 0.0027,$$

respectively, corresponding to energies of  $3.6 \times 10^6$  and  $2.5 \times 10^6$  e-volts. The neutron would receive three-quarters of this energy, i.e.,  $2.7 \times 10^6$ , or  $1.9 \times 10^6$  e-volts. Either of these two values are in good accord with the

\* 'Proc. Roy. Soc.,' A, vol. 141, p. 259 (1933).

† 'Nature,' vol. 133, p. 554 (1934).

‡ 'Nature,' vol. 133, p. 377 (1934).

§ Othman, Kinsey, and Rutherford, 'Proc. Roy. Soc.,' A, vol. 141, p. 722 (1933).

approximate value of  $2 \times 10^6$  e-volts found from our experiments, but Dee\* has now obtained expansion chamber photographs of the recoil nuclei, analysis of which suggests that the lower of the two figures is more nearly correct, and that the neutrons emitted at right angles to the bombarding beam of dipions are homogeneous in velocity, as required by our reaction

The recoiling  ${}^3_2\text{He}^3$  nucleus possesses an energy of about  $0.7 \times 10^6$  e-volts, i.e., a range of 5-6 mm as a maximum.† The thinnest mica window we have been able to use has a stopping power of 3 mm of air, and the residual range of 2-3 mm. is not sufficient for the particle to enter our counting chamber and produce a deflection of appreciable size. We have searched very carefully for such a doubly charged particle, both with a special type of counting chamber of small depth, and by looking for scintillations produced on a screen of zinc sulphide placed inside the apparatus itself and covered with aluminium of 2 mm. stopping power to prevent light and scattered dipions from reaching it. In both cases the presence of a very intense radiation which is strongly absorbed in a fraction of a centimetre of air, gave rise to so much disturbance as to render counting impossible. Thus, while we have not yet detected the  ${}^3_2\text{He}^3$  particles which we believe to be present, we have not yet obtained any evidence that they do not exist.

No evidence of the existence of an  ${}^3_2\text{He}^3$  isotope has been obtained by ordinary methods, although the possibility of its existence has been suggested at various times. It is not unlikely that while the new isotope may prove to be unstable over long periods it may yet have a sufficiently long life to be detected by counting methods and in the expansion chamber. We have not detected any after-effects lasting for a few seconds or more, suggesting the expulsion of a positive electron or other charged particle. If the  ${}^3_2\text{He}^3$  nucleus is unstable, there are a number of possibilities as to the mode of transformation.

It is evident that the experiments we have described suggest very strongly that the neutrons resulting from the bombardment of diplogen with diplogen are homogeneous in velocity, and since large yields are obtainable at comparatively low bombarding potentials they should serve as an almost ideal group for experimental work on the properties of neutrons.

In addition to a study of these transformations, we have made a number of observations on the neutrons, protons, and also  $\alpha$ -particles emitted from

\* Not yet published.

† The energy and range of the particle depends to some extent on the direction of emission relative to the bombarding particle, and will be greater in the forward direction.

lithium, beryllium, carbon, and other elements under dipion bombardment. We hope to give an account of these experiments in a subsequent paper.

In conclusion, we have to express our thanks for help with some of the experiments to Mr. Kempton and Mr. Westcott, and we acknowledge our indebtedness to Mr. G. R. Crowe for his technical assistance throughout. Dr. P. Hartek is indebted to the Rockefeller Foundation for a grant.

### *Summary*

An account is given of the effects observed when dipions are used to bombard targets of compounds containing heavy hydrogen. It is found that a group of protons of 14.3 cm. range is emitted in very large numbers. A shorter 1.6 cm. range group of singly charged particles is also observed, and it is shown that the two groups contain equal numbers of particles. A discussion of the reaction which gives rise to them is given, and reasons are advanced for supposing that the short-range group consists of nuclei of a new isotope of hydrogen of mass 3.0151. The number of particles emitted has been investigated as a function of the energy of the bombarding dipions, and the absolute yield for a pure dipion beam hitting a pure diplogen target is estimated to be about  $1 \text{ in } 10^6$  at 100,000 volts.

Neutrons have been observed in large numbers as a result of the same bombardment. It is shown that the energy of the neutrons is about  $2 \times 10^6$  e-volts, and it is suggested that they arise from an alternative mode of breaking up of the unstable form of helium nucleus formed initially by the union of two dipions. This other mode results in the expulsion of a neutron and a helium isotope of mass 3 in directions opposite to one another. If we calculate the mass of  ${}^3\text{He}$  from energy and momentum considerations of the ranges of the short-range groups emitted from  ${}^6\text{Li}$  when bombarded by protons, the energy of the neutron can be deduced and agrees well with experiment.

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*Experiments with High Velocity Positive Ions III—The Disintegration of Lithium, Boron, and Carbon by Heavy Hydrogen Ions*

By J D COCKCROFT, Ph.D, and E T S WALTON, Ph D

(Communicated by Lord Rutherford, O M, F R S—Received April 4, 1934)

1 *Introduction*

An important enlargement of the field of nuclear research has been effected by the experiments of Lawrence, Lewis, and Livingstone,\* who first used the ions of the heavy isotope of hydrogen to produce nuclear disintegrations. Their pioneer experiments showed that these ions, which will in future be termed "dipions,"† produced disintegrations of considerably greater complexity than those produced by protons of the same energies. Thus, Lawrence, Lewis, and Livingstone reported that from most of the substances bombarded, one or more groups of protons were emitted, whilst a group having a range of about 18 cm appeared to be emitted from every element investigated. This surprising result led these workers to the hypothesis that the "dipion" is unstable in a strong nuclear field and that it breaks up into a proton and a neutron with a liberation of energy of the order of 5 million volts, the neutron mass being taken to be 1.0006‡ to explain the observed proton energies.

In addition to the emission of long-range protons, Lawrence, Lewis, and Livingstone (*loc cit*) reported the emission of  $\alpha$ -particles from lithium, beryllium, boron, nitrogen, magnesium, and aluminium. The experiments of Rutherford, Oliphant, and Kinsey§ and the Wilson chamber photographs of Dee and Walton|| showed that the 13.2 cm.  $\alpha$ -particles from lithium originated from the disintegration of  $\text{Li}^6$  into two  $\alpha$ -particles, and suggested that  $\text{Li}^7$  disintegrated into two  $\alpha$ -particles and a neutron. No detailed studies have been made of the disintegration of the other elements.

The experiments described below have been made with a view to obtaining more evidence on the nature of the disintegrations in which protons are emitted, we have also studied the emission of  $\alpha$ -particles from boron under dipion bombardment.

\* 'Phys. Rev.', vol. 44, p. 55 (1933).

† 'Proc. Roy. Soc.', A, vol. 144, p. 1 (1934).

‡ 'Phys. Rev.', vol. 44, p. 778 (1933).

§ 'Proc. Roy. Soc.', A, vol. 141, p. 722 (1933).

|| 'Proc. Roy. Soc.', A, vol. 141, p. 733 (1933).

## 2 Apparatus

In these experiments, we have used the high voltage D C generator described in our previous paper,\* which gives a maximum steady potential of 700 kv. The tube for the acceleration of ions there described has been modified to give larger intensities of ions and more reliable operation. Porcelain cylinders of 12-inch diameter and 3 feet 6 inches length have been substituted for the glass cylinders used previously, a discharge tube of the type described by Rutherford and Oliphant† has been transferred from the apparatus of Dee and Walton to the new apparatus and the evacuation of the tube is now carried out by a Metropolitan-Vickers 04 Apiezon Oil Diffusion Pump which has a speed of 1200 litres/sec. for hydrogen. This great increase in pumping speed has increased considerably the reliability of operation of the tube, the new discharge tube enables total currents of 200 microamperes to be obtained, whilst, in spite of using double stage acceleration, the geometry of the electrodes is now such as to focus about 20 per cent of the ions into a beam having a diameter of only 1 cm. in the experimental chamber, thereby increasing considerably the fraction of disintegration particles which can be made to enter the recording apparatus.

## 3 The Detection of High Speed Protons

In most experiments we have used the scintillation method for fixing approximately the ranges of the particles emitted, carrying out a detailed study with the ionization chamber, linear amplifier, and oscillograph previously described. In interpreting records of proton emission, difficulty arises when the background disturbance of the oscillograph record due to electrical and acoustical "pick-up" is comparable with the oscillograph deflection due to the protons. The greatest care has been taken to reduce as far as possible the magnitude of this pick up, but in practice there is often a disturbance due to the discharge tube or to soft radiation which gives a background oscillation on the oscillograph. Consequently it is often necessary to count only deflections above some arbitrary size corresponding to protons approaching the end of their range. The absorption curves obtained are thus of a differential type, maxima being obtained near the end of the range of a group of protons.

In the present experiments we have not applied magnetic analysis to separate out the protons and dipions in our beams. Since we usually have a mixture of hydrogen and dipion in our discharge tube, the beam of ions contains

\* 'Proc Roy Soc,' A, vol 136, p. 619 (1932)

† 'Proc Roy Soc,' A, vol. 141, p. 259 (1933).

protons of full and half energy and dipions of full and two-thirds energy. This spread of the energy causes the peaks in the absorption curves to be less sharply defined, but does not prevent an accurate deduction of the range from being made.

In most of these experiments, an ionization chamber having a depth of 1 cm. has been used, in general a correction of 0.5 cm. has been made to the ranges obtained from the absorption curves to allow for this depth.

#### 4. *The Disintegration of Lithium*

A target of pure Kahlbaum lithium, which on exposure becomes covered with a layer of lithium hydroxide, was bombarded with a mixed beam of ions obtained from hydrogen containing approximately 10 per cent of diplogen. A target of 5 cm. diameter was used, the beam covering a diameter of about 1 cm., the distance from centre of target to recorder was 5 cm., and the solid angle subtended by the counter aperture at the recorder was about  $1/30$ . With this arrangement, the emission of particles having a range of about 30 cm. was observed, and the absorption curve of fig. 1 was obtained for an accelerating

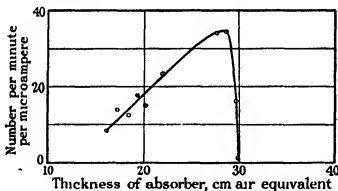


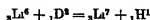
FIG. 1.—Protons from lithium

potential of 500 kv. The maximum number of particles counted was about 700 per minute. The observations show that a group of protons having a range of  $30.5 \text{ cm.} \pm 1 \text{ cm.}$  for an accelerating potential of 500 kv. is emitted from lithium. Observations were not carried out at absorptions less than 14 cm. owing to the presence of the strong  $\alpha$ -particle group of 13.2 cm. range. The range observed is consistent with the observation of Lawrence, Lewis, and Livingstone that protons of range up to 40 cm. were emitted from lithium fluoride when bombarded with their more energetic particles.



5 *The Mechanism of the Disintegration.*

If we make the assumption that the following nuclear reaction occurs



the kinetic energy change in the transmutation may be calculated from the mass change

$$\begin{aligned} & (6.0145 + 2.0136) - (7.0146 + 1.0078) \\ & = 0.0057 \pm 0.0007 = 5.3 \pm 0.6 \text{ million volts} \end{aligned}$$

Allowing for the kinetic energy of the deuterons and the energy taken by the recoiling nucleus, the disintegration protons may be expected to have an energy of  $5.0 \pm 0.5$  million volts. If we adopt Duncanson's figures\* for the range velocity function of the protons, protons of this energy would have a range  $35 \pm 5$  cm. The observed ranges would thus be consistent with the hypothesis as to the nature of the disintegration. If, on the other hand, we take the masses of  $\text{Li}^6$  and  $\text{Li}^7$  to be fixed by disintegration data, the expected energy in the protons is  $5.1 \pm 0.2$  million volts, which, from Duncanson's data would give a range of  $35 \pm 2$  cm. The agreement while fair is not as good as might be expected, the discrepancy may be due to the uncertainties in the range velocity relation for protons, to the emission of radiation in the transmutation, or it may be possible that the unstable nucleus  $\text{Be}^8$  may lose part of the energy imparted by the deuteron before breaking up.

Strong support for the proposed reaction is given by the experiments of Oliphant, Shire, and Crowther† on the transmutation of the separated isotopes of lithium, which showed that the 30 cm. protons result from the bombardment of  $\text{Li}^6$  by  $\text{H}^2$ .

6 *The Number of Disintegrations*

The absolute number of disintegrations cannot be obtained with any precision from our present experiments owing to the uncertainty in the composition of the beam. For the same reason, no accurate determination of the variation of the number of disintegrations with voltage has been made. The results for the mixed beam are, however, given in fig. 2. It is seen that there is no sharply defined lower limit to the voltage at which disintegrations can be observed, the curve being similar to that obtained for the 13.2 cm.  $\alpha$ -particles emitted in the disintegration



\* 'Proc Camb Phil Soc,' vol 30, p. 102 (1934)

† 'Nature,' vol. 133, p. 377, March 10 (1934).

We find that the number of 13.2 cm.  $\alpha$ -particles emitted is roughly twice the number of disintegration protons for dipion energies of 500 kv., so that the relative probability of the two types of disintegration is not very different.

### 7. The Disintegration of Carbon.

The absorption curves obtained for a target of Acheson graphite with the same experimental arrangement as for lithium are given in fig. 3, the maximum number of disintegration particles being of the same order as that found for

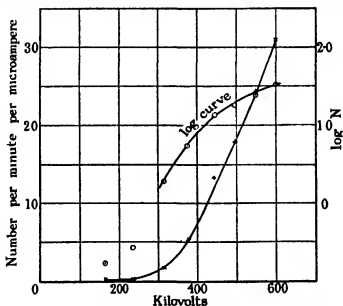


FIG. 2.—Protons from lithium, 27.8 cm. absorber

lithium. The curves suggest the presence of a group of particles having a range of  $14.0 \pm 1$  cm. for an accelerating potential of 500 kv. An analysis of the oscillograph deflections suggests that this group are protons, and it also indicates the presence of a weak proton group having a range of about 8 cm.

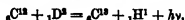
Taking the energy of the 14 cm. protons as 2.9 million volts, the kinetic energy change in the reaction is  $2.6 \pm 0.16$  million volts.

The range of the proton group obtained is about 1 cm. longer than would be expected from the observations of Lawrence, namely, that, using 1.2 million volt dipions, the proton range is 18 cm.

Since these experiments were completed, Lauritsen and Crane\* have reported the emission of a  $\gamma$ -ray having an energy of about 3 million volts when carbon

\* 'Phys. Rev.', vol. 45, p. 345 (1934).

is bombarded with deuterons. If we assume that the 14 cm. proton and the  $\gamma$ -ray are emitted at the same time from the reaction



we obtain the following value for the mass of  $\text{C}^{13}$

$$12.0036 + 2.0136 - 1.0078 - 0.0027 - 0.0032 = 13.0035 \pm 0.0006$$

The band spectrum mass for  $\text{C}^{13}$  is  $13.0039 \pm 0.0013$ . The mass obtained from the disintegration of  $\text{B}^{10}$  by  $\alpha$ -particles is  $13.0047 \pm 0.0005$  (Chadwick, Constable, and Pollard) \*

If the reaction assumed is correct, it might be expected that some transmutations would occur in which no  $\gamma$ -ray was emitted, the full energy appearing in the form of kinetic energy, so that protons of about 40 cm. range should be

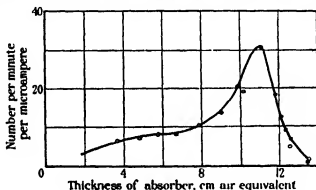


Fig. 3.—Protons from carbon. Dipion energy 500 kv

observed. It appears certain, however, that if they exist the number of such protons must be less than 1 in 500 of the number of 14 cm. protons.

On the other hand, the emission of the  $\gamma$ -ray observed by Lauritsen and Crane may be connected with the transformation of  $\text{C}^{13}$  to  $\text{N}^{13}$  under proton bombardment with the subsequent emission of a positive electron.† If this is so, then the mass of  $\text{C}^{13}$  calculated from both reactions would be about 13.006.

Evidence to be referred to later suggests that the 8 cm. proton group may be due to a layer of adsorbed oxygen.

Fig. 4 gives the observed variation in the number of disintegrations with voltage. These results were obtained from a beam of mixed ions, and an absolute determination of numbers has not yet been made.

\* 'Proc. Roy. Soc.,' A, vol. 130, p. 463 (1933).

† Cockcroft, Gilbert, and Walton, 'Nature,' vol. 133, p. 328 (1934).

8 *The Disintegration of Boron*

A study of the particles emitted from a target of boron under dipion bombardment has shown that very swift  $\alpha$ -particles are emitted together with at least three groups of protons. The absorption curve is given in fig 5.

By combining deductions from the shape of the absorption curve, fig 5, with an analysis of the distribution of the size of oscillograph deflections we find that both  $\alpha$ -particle and proton groups are present. A complete interpretation

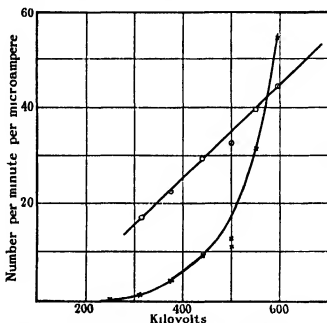


FIG. 4.—Dipions on carbon target, absorber 8.9 cm. of air

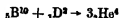
of these curves up to 16 cm. is difficult owing to the probable superposition of one or more proton groups on a continuous distribution of  $\alpha$ -particles. The existence of the following groups of particles seems, however, to be certain —

- (1)  $\alpha$ -particles of all ranges up to a maximum of  $15.0 \pm 1$  cm.
- (2) A proton group of range  $31 \pm 2$  cm.
- (3) A proton group of range  $58 \pm 2$  cm.
- (4) A proton group of range  $92 \pm 2$  cm.

The rise in the absorption curve from 6 to 10 cm. suggests the presence of another proton group, but until Wilson chamber photographs are available it is not possible to be certain of the interpretation.

The following tentative explanations are advanced for groups (1) to (4)

(1) The continuous distribution of  $\alpha$ -particles corresponds to the disintegration of  $B^{10}$  into three  $\alpha$ -particles



The absorption curve for this disintegration would be expected to be of a similar nature to that observed for the disintegration of  $B^{11}$  by protons\*. If

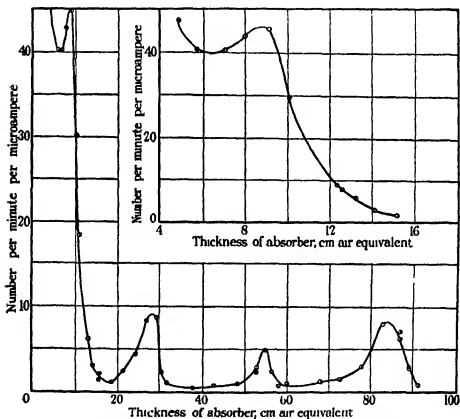


FIG 5 — $\alpha$ -particles and protons from boron Dipion energy 500 kv

the hump in the curve at 9 cm. is due to a superposed homogeneous group of protons, then the residual curve would have a shape similar to the  $B^{11}$  curve

If the end-point of the  $\alpha$ -particle curve corresponds to a disintegration in which one  $\alpha$ -particle is ejected in a direction opposite to the other two, then

\* Rutherford and Oliphant, 'Proc. Roy Soc.', A, vol. 141, p 259 (1933)

this  $\alpha$ -particle possesses two-thirds of the total kinetic energy. The total kinetic energy in the disintegration particles would thus be

$$(12.3 \pm 0.30) \times 3/2 \text{ million volts} = 18.45 \pm 0.45 \text{ million volts.}$$

If the mass of  $B^{10}$  be taken as  $10.0135 \pm 0.0005$  the expected kinetic energy is  $19.4 \pm 0.6$  million volts. The agreement is satisfactory, but it is very desirable to have further evidence about this disintegration from Wilson chamber photographs. It should be noted that some of the  $\alpha$ -particles from this reaction are swifter than any which have previously been discovered. The fastest  $\alpha$ -particles from radioactive bodies have a range of 11.5 cm. whilst the  $\alpha$ -particles discovered by Lawrence in the reaction  $Li^6 + D$  had a range of 14 cm.

(2), (3), and (4) assuming Duncanson's range velocity relation for protons, these groups correspond to protons of energy 4.7, 6.7, and 8.8 million volts, and could be explained by the following reaction.



If the proton of range 92 cm is emitted when all the energy of the reaction goes into kinetic energy, we may calculate the difference of mass of  $B^{10}$  and  $B^{11}$  as follows

$$\begin{aligned} B^{11} - B^{10} &= 2.0136 - 1.0078 + 0.0005 \text{ (K.E. of } D^2) \\ &\quad - 0.0102 \text{ (K.E. after disintegration)} \\ &= 0.9961 \pm 0.0004 \end{aligned}$$

The difference of mass from Aston's determination is  $11.0110 - 10.0135 = 0.9975 \pm 0.0007$ . Thus the agreement is not very good. If, on the other hand, we were to assume that the masses of  $B^{10}$  and  $B^{11}$  could be obtained from the end-point of the continuous  $\alpha$ -particle distributions in the transmutations by deuterons and protons respectively, we should obtain very close agreement with these experiments.

The two groups of lower energy may correspond to disintegrations in which energy of 2 and 4 million volts respectively is emitted in the form of  $\gamma$ -radiation.

The total number of particles in groups (2), (3), and (4) is approximately one-third of the maximum number of  $\alpha$ -particles observed. This suggests that the disintegration leading to three  $\alpha$ -particles has about the same probability as the disintegration leading to  $B^{11}$ .

*9 The Emission of Particles on the Bombardment of Heavy Elements  
by Diprotons*

A detailed study has been made of the particles emitted from copper, copper oxide, iron, iron oxide, tungsten, and silver. In order to facilitate comparison of these elements the target illustrated by fig 6 was constructed so that four targets could be interchanged by rotation of a ground joint. With this arrangement the solid angle subtended by the window at the target was increased to 0.25, a still further increase in the number of particles observed was obtained by using a helium-diplogen mixture containing 30% of diplogen. A series of eight experiments was carried out, using diplogen energies of 500 kv.

(i) and (ii) The emission of particles from carbon, copper, copper oxide, copper, and iron was compared in one series of experiments and of tungsten, silver, iron oxide, and iron in another, the absorption curves of figs 7, *a*, and 7, *b*, being obtained. Observations could not be carried out on silver at less absorber than 6 cm. owing to the emission of soft radiation in considerable intensity.

From these curves and from an analysis of the size of the oscillograph deflections we conclude—

- (a) that a group of protons of range about 14 cm. is emitted from all these targets, the numbers from carbon being about ten times the numbers from other targets,
- (b) that a group of protons of range 8 cm. is emitted from all the targets,
- (c) that a group of singly charged particles of range about 4 cm. is common to all targets except carbon,
- (d) that  $\alpha$ -particles are emitted from all the targets, the numbers falling rapidly for absorbers greater than 2 cm., a few particles being observed up to 5 cm.
- (e) that whilst the 8 cm. group is stronger from  $\text{Fe}_2\text{O}_3$  than from Fe, the relative strength of the 14 cm. groups are reversed.

(iii) A strip of the tungsten used in experiment (ii) was washed in succession in caustic soda, distilled water, ether, caustic soda, and distilled water and

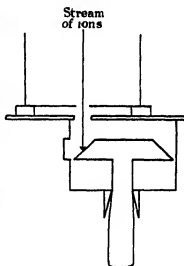


FIG 6

then bombarded. The absorption curve of fig 8, *b*, was then obtained, showing that whereas the short range group of particles and the number of particles at 8 cm. absorber had roughly the same relative magnitude as in experiment (u) the 14 cm. peak was reduced in numbers to about one-third of the 8 cm.

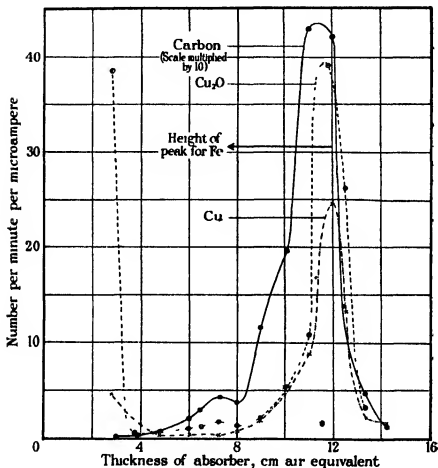


FIG. 7, a.

group. In addition the residual group appears now to have a rather longer range than that of the original group.

(iv) At the suggestion of Dr. Oliphant the target was heated in position to about  $2000^\circ\text{C}$  by passing a heavy current through it, and the emission of particles observed under these conditions and immediately after heating. The results showed a considerable diminution in the number of particles in the 8 cm. groups and short-range groups, whilst the numbers in the 14 cm.



group were reduced by a further factor of 2, the curve obtained after heating being shown on fig 8, c

(v) The target was left for 18 hours in the apparatus and then bombarded whilst cold. The number of particles observed was if anything smaller than immediately after heating the target, showing that no new layers of impurities had been adsorbed during this period

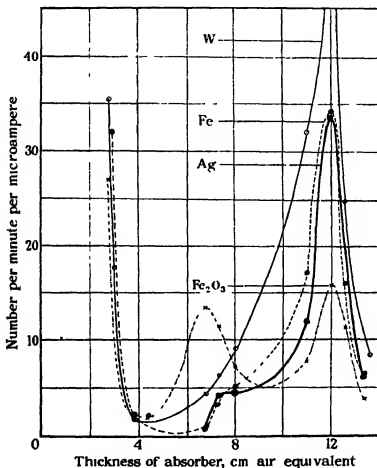


FIG 7, b—Diplon energy 500 kv

(vi) Carbon dioxide was admitted to the apparatus to a pressure of 1 cm, the target flashed at about 1600° C and again bombarded. No change in the number of particles was observed from experiment (v), showing that no surface layer had been adsorbed in sufficient amount to give rise to an appreciable effect.

(vii) After being cleaned by heating, the target was taken out and oxidised by passing an electric current through it, replaced and again bombarded. The absorption curve obtained is shown in fig. 8, *d*. The short-range group and

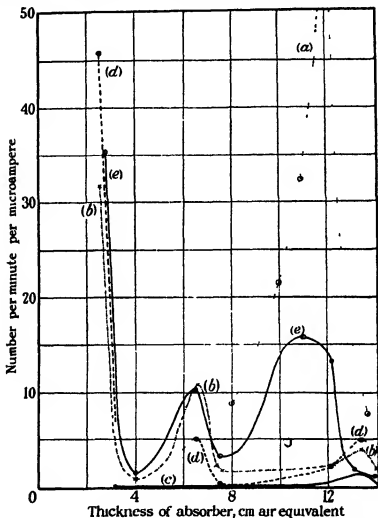


FIG. 8.—Diplon energy 500 kv (a) Original tungsten, (b) chemically cleaned tungsten, (c) after heating, (d) after oxidizing, (e) after reheating.

8 cm. group have again appeared strongly together with a few particles at absorptions of 14 cm. about equal in numbers to those obtained in (iii)

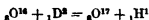
(viii) The target was cleaned by heating in vacuum, was removed and oxidized and then handled in order to reproduce the treatment which a metal

might reasonably receive before mounting in an apparatus. The target was then replaced and bombarded and the curves of fig. 8, *e*, were obtained. We see that all three peaks now appear strongly, the long-range peak is again stronger than the 8 cm. peak and is now restored to its former position.

These experiments may be summarized as follows —

(1) The 8 cm. proton group, the 4 cm. proton group, and the short-range  $\alpha$ -particle groups disappear when the target is heated strongly and reappear when the target is oxidized. The evidence suggests therefore that these groups are due to oxygen.

Other experiments carried out by Rutherford and Oliphant\* have led to similar conclusions and they propose the following explanation for the reaction leading to the 8 cm. protons



the observed range being in good accord with the accepted masses for  $\text{O}^{16}$  and  $\text{O}^{17}$ . The explanation of the 4 cm. proton group must await further experiments, the energy difference between the two groups is 0.75 million volts.

(2) The long-range proton group from the heavy elements appears to be complex, the greater part of the effect being removed by chemical cleaning of the surfaces, a small residual effect remaining due to protons having a slightly longer range. The coincidence of the range of the majority of the protons with the long-range group from carbon suggests therefore that a thin film of grease or other carbon impurity is responsible for the greater part of the effects.

An alternative explanation seemed, however, to be offered by the experiments of Rutherford, Harteck, and Oliphant,† which showed that on bombarding by deuterons, ammonium sulphate, in which the ordinary hydrogen had been largely replaced by deuterium, very great numbers of particles were emitted, half of the particles having a range of 1.7 cm. and half a range of about 1.4 cm. It seemed, therefore, possible that the whole of our results might be explained by deuterium being absorbed on carbon from the bombarding stream, the adsorbed layer being then disintegrated by the deuteron stream.

In order to test this hypothesis, we have carefully compared the ranges of the proton groups emitted from  $(\text{ND}_4)_2\text{SO}_4$  and carbon under identical conditions. If all the results are due to adsorbed deuterium, we ought to get the

\* *In course of publication.*

† 'Nature,' vol. 133, p. 413, March 17 (1934).

same 14 cm. range from the two substances, and we ought to observe large numbers of singly charged particles having a range of about 2 cm. from carbon.

The results of these experiments are given by figs. 9 and 10. In fig. 9 the absorption curves for the two substances are given, showing that the range of the protons from  $(\text{ND}_4)_2\text{SO}_4$  is greater by about 1 cm. than the range of the protons from carbon. A still more conclusive proof of the difference between the two reactions is given by fig. 10 in which we plot the distribution in size of the oscillograph deflections for the two substances for small absorptions. Remembering that a large deflection corresponds to a proton near the end of

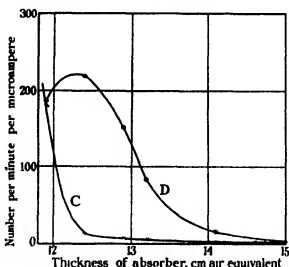


FIG. 9.—Comparison of ranges of long range protons from  $(\text{ND}_4)_2\text{SO}_4$  and C. D accelerating voltage of dipions 150 kv, proton range  $14 \pm 0.5$  cm. C accelerating voltage of dipions 370 kv, proton range  $12.5 \pm 0.5$  cm.

its range, we see at once that there are very few short-range particles from carbon compared with the number from  $(\text{ND}_4)_2\text{SO}_4$ . Experiments with greater thickness of absorber showed that the number of 14 cm. protons are of the same order of magnitude for the two targets.

We conclude that—

- (1) most of the 14 cm. protons emitted from carbon are due to the transmutation of  $\text{C}^{12}$  into  $\text{C}^{13}$ ,
- (2) there is probably a small additional group of protons of range 15 cm. at 500 kv due to the adsorption of diplogen and the subsequent disintegration of the adsorbed layer,

- (3) the greater part of the effects observed with the heavy elements is due to thin films of grease or other form of carbon impurity adsorbed on the surface ,
- (4) a smaller number of 15 cm protons are emitted from the heavy elements due to the adsorption of diplogen from the bombarding stream

## 10 Conclusions

We may summarize the experimental results by stating (i) that the emission of long-range protons is in all substances investigated (D, Li, B, C, O) bound up

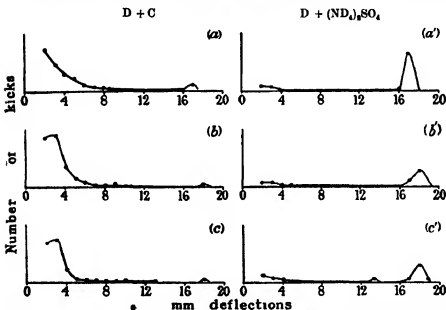


FIG 10—Distribution in size of oscillograph deflections for C + D and  $(\text{ND}_4)_2\text{SO}_4$  + D

- (a) and (a') 0.63 cm absorber  
 (b) and (b') 1.23 cm absorber  
 (c) and (c') 2 cm absorber

with the transformation of an element into the next heavier isotope of that element, the neutron being captured and the proton ejected, (ii) that when transmutations leading to the emission of an  $\alpha$ -particle are possible (Li, B) the probability of this transmutation is of the same order as that of the first type, (iii) that the emission of protons from the heavier elements (Cu, Fe, Ag, W) is in our experiments due to films of impurities of oxygen, carbon, and adsorbed diplogen.

We wish finally to express our appreciation of Lord Rutherford's constant interest and encouragement, and to thank Dr Harteck for his work in the preparation of the diplogen. We are indebted to the Government Grant Committee of the Royal Society for a grant in aid of the purchase of the fast pumping equipment used, one of us (E T S W) wishes to acknowledge the grant of a Senior Research Award of the Department of Scientific and Industrial Research

### *Summary*

The disintegration of lithium, boron, and carbon, when bombarded by fast ions of heavy hydrogen, has been examined. For lithium, a group of protons of 30.5 cm. range has been observed, this can be interpreted as resulting from the transformation of  $\text{Li}^6$  to  $\text{Li}^7$ . Boron gives proton groups of ranges 92 cm., 58 cm., and 31 cm. together with a continuous distribution of  $\alpha$ -particles with ranges up to a maximum of 15 cm. Carbon gives a group of protons with a range of 14 cm. These would correspond with the 18 cm. group observed by Lawrence from all targets if allowance is made for the higher energies of his bombarding particles.

A 14 cm. group of protons has been observed from a number of heavy elements. Experiments have been carried out which indicate that these are probably due to carbon present on the surface in a thin film of grease. A smaller number of protons having a slightly longer range are also emitted, these are probably produced from the disintegration of diplogen adsorbed on the surface. Layers of oxygen on the surface may also give rise to spurious effects.

The emission of a fast group of protons always appears to be connected with the change of one isotope of an element into its next heavier isotope.

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